PHENYLALANINE DERIVATIVES

This invention relates to a series of phenylalanine derivatives, to compositions containing them, to processes for their preparation, and to their use in medicine.

Over the last few years it has become increasingly clear that the physical interaction of inflammatory leukocytes with each other and other cells of the body plays an important role in regulating immune and inflammatory responses [Springer, T A. Nature, <u>346</u>, 425, (1990); Springer, T. A. Cell <u>76</u>, 301, (1994)]. Many of these interactions are mediated by specific cell surface molecules collectively referred to as cell adhesion molecules.

The adhesion molecules have been sub-divided into different groups on the basis of their structure. One family of adhesion molecules which is believed to play a particularly important role in regulating immune and inflammatory responses is the integrin family. This family of cell surface glycoproteins has a typical non-covalently linked heterodimer structure. At least 14 different integrin alpha chains and 8 different integrin beta chains have been identified [Sonnenberg, A. Current Topics in Microbiology and immunology, 184, 7, (1993)]. The members of the family are typically named according to their heterodimer composition although trivial nomenclature is widespread in this field. Thus the integrin termed $\alpha 4\beta 1$ consists of the integrin alpha 4 chain associated with the integrin beta 1 chain, but is also widely referred to as Very Late Antigen 4 or VLA4. Not all of the potential pairings of integrin alpha and beta chains have yet been observed in nature and the integrin family has been subdivided into a number of subgroups based on the pairings that have been recognised [Sonnenberg, A. ibid].

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The importance of cell adhesion molecules in human leukocyte function has been further highlighted by a genetic deficiency disease called Leukocyte Adhesion Deficiency (LAD) in which one of the families of leukocyte integrins is not expressed (Marlin, S. D. <u>et al</u> J. Exp. Med. 164. 855 (1986)]. Patients with this disease have a reduced ability to recruit

leukocytes to inflammatory sites and suffer recurrent infections which in extreme cases may be fatal.

The potential to modify adhesion molecule function in such a way as to beneficially modulate immune and inflammatory responses has been extensively investigated in animal models using specific monoclonal antibodies that block various functions of these molecules [e.g. Issekutz, T. B. J. Immunol. 3394, (1992); Li, Z. <u>et al</u> Am. J. Physiol. 263, L723, (1992); Binns, R. M. <u>et al</u> J. Immunol. 157, 4094, (1996)]. A number of monoclonal antibodies which block adhesion molecule function are currently being investigated for their therapeutic potential in human disease.

One particular integrin subgroup of interest involves the a4 chain which can pair with two different beta chains \$1 and \$7 (Sonnenberg, A. ibid). The a461 pairing occurs on many circulating leukocytes (for example lymphocytes, monocytes and eosinophils) although it is absent or only present at low levels on circulating neutrophils. 0481 binds to an adhesion molecule (Vascular Cell Adhesion Molecule-1 also known as VCAM-1) frequently up-regulated on endothelial cells at sites of inflammation [Osborne, L. Cell, 62, 3, (1990)]. The molecule has also been shown to bind to at least three sites in the matrix molecule fibronectin [Humphries, M. J. et al. Ciba Foundation Symposium, 189, 177, (1995)]. Based on data obtained with monoclonal antibodies in animal models it is believed that the interaction between $\alpha 4\beta 1$ and ligands on other cells and the extracellular matrix plays an important role in leukocyte migration and activation [Yednock, T. A. et al, Nature, 356, 63, (1992); Podolsky, D. K. et al. J. Clin, Invest, 92, 373, (1993); Abraham, W. M. et al. J. Clin, Invest. 93, 776, (1994)].

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The integrin generated by the pairing of α4 and β7 has been termed LPAM-1 [Holzmann, B and Weissman, I. EMBO J. §, 1735, (1989)] and like α4β1, binds to VCAM-1 and fibronectin. In addition, α4β7 binds to an adhesion molecule believed to be involved in the homing of leukocytes a mucosal tissue termed MAdCAM-1 [Berlin, C. et al., Cell, Z4, 185, (1993)]. The interaction between α4β7 and MAdCAM-1 may also be important at

sites of inflammation outside of mucosal tissue [Yang, X-D. <u>et al</u>, PNAS, 91, 12604 (1994)].

Regions of the peptide sequence recognised by α4β1 and α4β7 when they bind to their ligands have been identified. α4β1 seems to recognise LDV, IDA or REDV peptide sequences in fibronectin and a QIDSP sequence in VCAM-1 [Humphries, M. J. et al, ibid] whilst α4β7 recognises a LDT sequence in MAdCAM-1 [Briskin, M. J. et al, J. Immunol. 156, 719, (1996)]. There have been several reports of inhibitors of these interactions being designed from modifications of these short peptide sequences (Cardarelli, P. M. et al, J. Biol. Chem. 269, 18668, (1994); Shroff, H. N. Bioorganic. Med. Chem. Lett. 6, 2495, (1996); Vanderslice, P. J. Immunol. 158, 1710, (1997)]. It has also been reported that a short peptide sequence derived from the α4β1 binding site in fibronectin can inhibit a contact hypersensitivity reaction in a trinitrochlorobenzene sensitised mouse [Ferguson, T. A. et al, PNAS 88, 8072, (1991)].

Since the alpha 4 subgroup of integrins are predominantly expressed on leukocytes their inhibition can be expected to be beneficial in a number of immune or inflammatory disease states. However, because of the ubiquitous distribution and wide range of functions performed by other members of the integrin family it is very important to be able to identify selective inhibitors of the alpha 4 subgroup.

25 We have now found a group of compounds which are potent and selective inhibitors of α4 integrins. Members of the group are able to inhibit α4 integrins such as α4β1 and/or α4β7 at concentrations at which they generally have no or minimal inhibitory action on α integrins of other subgroups. The compounds are thus of use in medicine, for example in the prophylaxis and treatment of immune or inflammatory disorders as described hereinafter.

Thus according to one aspect of the invention we provide a compound of formula (1)

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$$R^{1}(A|k^{1})_{s}(L^{1})_{s}$$

$$= A^{2} \qquad (A|k^{2})_{m}$$

$$= C(R^{2}) - N(R^{3})COHet$$

$$= R$$

wherein

R is a carboxylic acid or a derivative thereof:

5 R1 is a hydrogen atom or a hydroxyl, straight or branched alkoxy or optionally substituted cycloaliphatic, polycycloaliphatic, heterocycloaliphatic, polyheterocycloaliphatic, aromatic or heteroaromatic group; Alk1 is an optionally substituted aliphatic or heteroaliphatic chain;

L1 is a linker atom or group;

10 r and s, which may be the same or different, is each zero or an integer 1 provided that when r is zero R1 is an optionally substituted cycloaliphatic. polycycloaliphatic, heterocycloaliphatic, polyheterocycloaliphatic, aromatic or heteroaromatic group:

Ra and Rb, which may be the same or different is each an atom or group -L2(CH2)pL3(Rc)q in which L2 and L3 is each a covalent bond or a linker atom or group, p is zero or the integer 1, q is an integer 1, 2 or 3 and Rc is a hydrogen or halogen atom or a group selected from straight or branched alkyl, -ORd (where Rd is a hydrogen atom or an optionally substituted straight or branched alkyl groupl, -SRd, -NRdRe, [where Re is as just defined for Rd and may be the same or different], -NO2,-CN, -CO2Rd, -SO₂H. -SO₂Rd. -OCO₂Rd. -CONRdRd. -OCONRdRd. -CSNRdRd. -CORd. -N(Rd)CORe, N(Rd)CSRe, -SO2N(Rd)(Re), -N(Rd)SO2Re, -N(Rd)CONReRf (where Rf is a hydrogen atom or an optionally substituted straight or branched alkyl group], -N(Rd)CSNReRf or -N(Rd)SO2NReRf;

25 Alk2 is a straight or branched alkylene chain;

m is zero or an integer 1;

R2 is a hydrogen atom or a methyl group:

R3 is a hydrogen atom or a straight or branched alkyl group;

Het is an optionally substituted heteroaromatic group; and the saits, solvates, hydrates and N-oxides thereof. It will be appreciated that compounds of formula (1) may have one or more chiral centres. Where one or more chiral centres is present, enantiomers or diastereomers may exist, and the invention is to be understood to extend to all such enantiomers, diasteromers and mixtures thereof. including racemates. Formula (1) and the formulae hereinafter are intended to represent all individual isomers and mixtures thereof, unless stated or shown otherwise

In the compounds of formula (1), derivatives of the carboxylic acid group R include carboxylic acid esters and amides. Particular estes and amides 10 include -CO2Alk4 and -CON(R4)2 groups as described herein.

When in the compounds of the invention L1 is present as a linker atom or group it may be any divalent linking atom or group. Particular examples include -O- or -S- atoms or -C(O)-, -C(O)O-, -C(S)-, -S(O)-, -S(O)2-, -N(R4)-[where R4 is a hydrogen atom or a straight or branched alkyl group]. -CON(R4)-, -OC(O)N(R4)-, -CSN(R4)-, -N(R4)CO-, -N(R4)C(O)O-, -N(R4)CS-, -S(O)N(R4)-, -S(O)2N(R4)-, -N(R4)S(O)-, -N(R4)S(O)2-. -N(R4)CON(R4)-. -N(R4)CSN(R4)-, -N(R4)SON(R4)- or -N(R4)SO₂N(R4)groups. Where the linker group contains two R4 substituents, these may 20 be the same or different.

Alk2 in the compounds of the invention may be for example a straight or branched C1-3alkylene chain. Particular examples include -CH2-, -CH(CH3)-, -C(CH3)2- and -(CH2)2-.

When R3 and/or R4 in the compounds of formula (1) is a straight or branched alkyl group it may be a straight or branched C1-salkyl group, e.g. a C1.3alkyl group such as a methyl or ethyl group.

When Alk1 in compounds of formula (1) is an optionally substituted aliphatic chain it may be an optionally substituted C1-10 aliphatic chain. Particular examples include optionally substituted straight or branched chain C1-6 alkyl, C2-6 alkenyl, or C2-8 alkynyl chains.

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Heteroaliphatic chains represented by Alk¹ include the aliphatic chains just described but with each chain additionally containing one, two, three or four heteroatoms or heteroatom-containing groups. Particular heteroatoms or groups include atoms or groups L⁴ where L⁴ is as defined above for L¹ when L¹ is a linker atom or group. Each L⁴ atom or group may interrupt the aliphatic chain, or may be positioned at its terminal carbon atom to connect the chain to the atom or group R¹.

Particular examples of aliphatic chains represented by Alk1 include optionally substituted -CH2-, -CH2CH2-, -CH(CH3)-, -C(CH3)2-, -(CH₂)₂CH₂-, -CH(CH₃)CH₂-, -(CH₂)₃CH₂-, -CH(CH₃)CH₂CH₂-, -CHoCH(CHa)CHo-, -C(CHa)oCHo-, -(CHo)aCHo-, -(CHo)aCHo-, -CHCH-, -CH2CHCH-, -CHCHCH2CH2-, -CH₂CHCHCH₂-. -(CH2)2CHCH-, -CC-, -CCCH2-, -CH2CC-, -CCCH2CH2-, -CH2CCCH2-, or -(CH₂)₂CC- chains. Where appropriate each of said chains may be optionally interrupted by one or two atoms and/or groups L4 to form an optionally substituted heteroaliphatic chain. Particular examples include optionally substituted -L4CH2-, -CH2L4CH2-, -L4(CH2)2-, -CH2L4(CH2)2-, - (CH₂)₂L⁴CH₂-, -L⁴(CH₂)₃- and -(CH₂)₂L⁴(CH₂)₂- chains. The optional substituents which may be present on aliphatic or heteroaliphatic chains represented by Alk1 include one, two, three or more substituents selected from halogen atoms, e.g. fluorine, chlorine, bromine or iodine atoms, or hydroxyl, C₁₋₆aikoxy, e.tg, methoxy or ethoxy, thiol, C₁₋₆aikylthio e.g. methylthic or ethylthic, amino or substituted amino groupss. Substituted amino groups include -NHR4 and -N(R4)2 groups where R4 is a straight or branched alkyl group as defined above. Where two R4 groups are present these may be the same or different. Particular examples of substituted chains represented by Alk1 include those specific chains just described substituted by one, two, or three halogen atoms such as fluorine atoms, for example chains of the type -CH(CF₃)-, -C(CF₃)₂- -CH₂CH(CF₃)-, -CH₂C(CF₃)₂-, -CH(CF₃)- and -C(CF₃)₂CH₂.

Alkoxy groups represented by R^1 in compounds of the invention include straight of branched C_{1-6} alkoxy groups such as methoxy and ethoxy groups.

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When R1 is present in compounds of formula (1) as an optionally substituted cycloaliphatic group it may be an optionally substituted C₃₋₁₀ cycloaliphatic group. Particular examples include optionally substituted C₃₋₁₀cycloalkyl, e.g. C₃₋₇cycloalkyl, C₃₋₁₀cycloalkenyl or C₃₋₁₀cycloalkenyl

Optionally substituted heterocycloaliphatic groups represented by R¹ include the optionally substituted cycloaliphatic groups just described for R¹ but with each group additionally containing one, two, three or four heteroatoms or heteroatom-containing groups L² as just defined.

Optionally substituted polycycloaliphatic groups represented by R¹ include optionally substituted C₇₋₁₀ bi- or tricycloalkyl or C₇₋₁₀bi- or tricycloalkenyl groups. Optionally substituted polyheterocycloaliphatic groups represented by R¹ include the optionally substituted polycycloalkyl groups just described, but with each group additionally containing one, two, three or four L² atoms or groups.

Particular examples of R1 cycloaliphatic, polycycloaliphatic, heterocycloaliphatic and polyheterocycloaliphatic groups include optionally substituted cyclopropyl, cyclobetyl, cyclopentyl, cyclohexyl, cyclohexyl, 2-cyclopenten-1-yl, adamantyl, norborneyl, norbornenyl, pyrroline, e.g. 2- or 3-pyrrolinyl, pyrrolidinyl, pyrrolidinone, oxazolidinyl, oxazolidinone, dioxotanyl, e.g. 1,3-dioxotanyl, imidazolinyl, e.g. 2-imidazolinyl, imidazolidinyl, pyrazolinyl, e.g. 2-pyrazolinyl, pyrazolidinyl, thiazolidinyl, thiazolidinyl, pyrayl, e.g. 2- or 4-pyranyl, piperidinyl, piperidinone, 1,4-dioxanyl, morpholinyl, morpholinone, 1,4-dithianyl, thiomorpholinyl, piperazinyl, 1,3,5-trithianyl, oxazinyl, e.g. 2H-1,2- or 4H-1,4- oxazinyl, isoxazinyl, oxathlazinyl, e.g. 1,2,5 or 1,2,6-oxathlazinyl, or 1,3,5-oxadiazinyl groups.

The optional substituents which may be present on the R¹ cycloaliphatic, polycycloaliphatic, heterocycloaliphatic or polyheterocycloaliphatic groups include one, two, three or more substituents represented by R⁵ in which R⁵ is selected from halogen atoms, e.g. fluorine, chlorine, bromine or iodine atoms, or C₁-galkyl, e.g. methyl or ethyl, haloC₁-galkyl, e.g.

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halomethyl or haloethyl such as difluoromethyl or trifluoromethyl, hydroxyl, C_{1-6} alkoxy, e.g. methoxy or ethoxy, halo C_{1-6} alkoxy, e.g. halomethoxy or haloethoxy such as difluoromethoxy or trifluoromethoxy, thiol, C_{1-6} alkylthio e.g. methylthio or ethylthio, $-N(R^4)_2$, -CN, $-CO_2R^4$, $-NO_2$, $-CON(R^4)_2$, $-CSN(R^4)_2$, $-COR^4$, $-CSN(R^4)_2$, $-N(R^4)COR^4$, $-N(R^4)CSR^4$, $-SO_2N(R^4)_2$, $-N(R^4)SO_2R^4$, $-N(R^4)CON(R^4)_2$, $-N(R^4)CSN(R^4)_2$ and $-N(R^4)SO_2N(R^4)_2$ groups. In these substituents the group R^4 when present is a hydrogen atom or a straight or branched alkyl group as defined above. Where more than one R^4 group is present in a substituent each group may be the same or different. The substituent may be present on any available carbon atom or where appropriate any nitrogen atom, in the R^1 group

In the compounds of formula (1), optionally substituted aromatic groups represented by the group R¹ include for example monocyclic or bicyclic fused ring Ce-12 aromatic groups, such as phenyl, 1- or 2-naphthyl, 1- or 2-tetrahydronaphthyl, indanyl or indenyl groups, optionally substituted by one, two, three or more -L²(CH₂)_pL³(R°)_q atoms or groups, where L², L³, p and q are as previously defined and R° is as previously defined but is other than a hydrogen atom when L² and L³ is each a covalent bond and p is zero.

Optionally substituted heteroaromatic groups, represented by the group R1 or Het in compounds of formula (1) include for example optionally substituted C1-9 heteroaromatic groups containing for example one, two, three or four heteroatoms selected from oxygen, sulphur or nitrogen atoms. In general, the heteroaromatic groups may be for example monocyclic or bicyclic fused ring heteroaromatic groups. Monocyclic heteroaromatic groups include for example five- or six-membered heteroaromatic groups containing one, two, three or four heteroatoms selected from oxygen, sulphur or nitrogen atoms. Bicyclic heteroaromatic groups include for example nine- to thirteen-membered fused-ring heteroaromatic groups containing one, two or more heteroatoms selected from oxygen, sulphur or nitrogen atoms.

Particular examples of heteroaromatic groups of these types include optionally substituted pyrrolyl, furyl, thienyl, imidazolyl, N-C₁₋₆aimidazolyl,

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oxazolyl, isoxazolyl, thiazolyl, isothiazolyl, pyrazolyl, 1,2,3-triazolyl, 1,2,4-1.2.3-oxadiazolyl, 1.2,4-oxadiazolyl, 1,2,5-oxadiazolyl, 1,3,4oxadiazolyl, 1,3,4-thiadiazole, pyridyl, pyrimidinyl, pyridazinyl, pyrazinyl, 1.3.5-triazinvl. 1.2.4-triazinyl, 1,2,3-triazinyl, benzofuryl, [2,3-dihydro]benzofuryl, benzothlenyl, benzotriazolyl, indolyl, isoindolyl, benzimidazolyl, imidazof1.2-alpvridyl, benzothiazolyl, benzoxazolyl, benzopyranyl, (3.4dihydro]benzopyranyl, quinazolinyl, naphthyridinyl, pyrido(3.4-b]pyridyl, pyrido[3.2-b]pyridyl, pyrido[4,3-b]pyridyl, quinolinyl, isoquinolinyl, tetrazolyl, 5,6,7,8-tetrahydroguinolinyl, 5,6,7,8-tetrahydroisoguinolinyl, and imidyl, e.g. succinimidyl, phthalimidyl, or naphthalimidyl such as 1,8naphthalimidyl.

Optional substituents which may be present on R1 heteroaromatic groups include one, two, three or more -L2(CH2)nL3(Rc)n atoms on groups as just defined.

Examples of the substituents represented by Ra and Rb in compounds of formula (1) and which may be present on aromatic or heteroaromatic groups represented by R1 include atoms or groups -L2(CH2)pLRc, -L2(CH2)nRc, -L2Rc, -(CH2)nRc and -Rc wherein L2, (CH2)n, L and Rc are as defined above. Particular examples of such substituents include -L2CH2L2R6, -L2CH(CH3)L3R6, -L2(CH2)2L3R6, -2CH2R6, -L2CH(CH3)R6, -L2(CH2)2Rc, -CH2Rc, -CH(CH3)Rc and -(CH2)2Rc grouns.

Thus each of Ra and Rb and, where present, substituents on R1 aromatic or heteroaromatic groups in compounds of the invention may be for example selected from a hydrogen atom, a halogen atom, e.g. a fluorine, chlorine, bromine or iodine atom, or a C1-Balkyl, e.g. methyl, ethyl, npropyl, i-propyl, n-butyl or t-butyl, C1-6alkylamino, e.g. methylamino or ethylamino, C₁₋₆hydroxyalkyl, e.g. hydroxymethyl, hydroxyethyl or -C(OH)(CF3)2, carboxyC1.salkvl, e.g. carboxyethyl, C1-salkylthio e.g. methylthio or ethylthio, carboxyC1-6alkylthio, e.g. carboxymethylthio, 2carboxyethylthio or 3-carboxypropylthio, C1-6alkoxy, e.g. methoxy or ethoxy, hydroxyC₁₋₆alkoxy, e.g. 2-hydroxyethoxy, haloC₁₋₆alkyl, e.g. -CF₃, 35 -CHF2, CH2F, haloC1-6alkoxy, e.g. -OCF3,-OCHF2,-OCH2F, C1-6alkylamino, e.g. methylamino or ethylamino, amino (-NH2), aminoC1-6alkyl, e.g.

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aminomethyl or aminoethyl, C1-sdialkylamino, e.g. dimethylamino or diethylamino, C1-salkylaminoC1-salkyl, e.g. ethylaminoethyl, C1-sdialkylaminoC1-salkyl, e.g. diethylaminoethyl, aminoC1-salkoxy, e.g. aminoethoxy, C1-salkylaminoC1-salkoxy, e.g. methylaminoethoxy, C1-sdialkylaminoC1-6alkoxy, e.g. dimethylaminoethoxy, diethylaminoethoxy, isopropylaminoethoxy, or dimethylaminopropoxy, nitro, cyano, amidino, hydroxyl (-OH), formyl [HC(O)-], carboxyl (-CO2H), -CO2R12, C1.6 alkanoyi e.g. acetyl, thiol (-SH), thioC1-6alkyl, e.g. thiomethyl or thioethyl, sulphonyl (-SO₃H). C_{1-salkylsulphonyl, e.g. methylsulphonyl,} aminosulphonyl (-SO₂NH₂), C₁₋₆alkylaminosulphonyl, e.g. methylaminosulphonyl or ethylaminosulphonyl, C1-sdialkylaminosulphonyl, e.g. dimethylaminosulphonyl or diethylaminosulphonyl, phenylaminosulphonyl, carboxamido (-CONH2). C1-6alkylaminocarbonyl, e.g. methylaminocarbonyl or ethylaminocarbonyl, C1-sdialkylaminocarbonyl, e.g. dimethylaminocarbonyl or diethylaminocarbonyl, aminoC_{1-s}alkylaminocarbonyl, e.g. aminoethylaminocarbonyl, C1-6dialkylaminoC1-6alkylaminocarbonyl, e.g. diethylaminoethylaminocarbonyl, aminocarbonylamino, C1-6alkylaminocarbonylamino, e.g. methylaminocarbonylamino or ethylaminocarbonylamino, C1-edialkylaminocarbonylamino, e.g. dimethylaminocarbonylamino or diethylaminocarbonylamino. C1-salkylaminocabonylC1. salkylamino, e.g. methylaminocarbonylmethylamino, aminothiocarbonylamino. C1-salkylaminothiocarbonylamino, e.g. methylaminothiocarbonylamino or ethylaminothiocarbonylamino. C1-edialkylaminothlocarbonylamino, e.g. dimethylaminothiocarbonylamino or diethylaminothiocarbonylamino, C1-salkylaminothiocarbonylC1-salkylamino, e.g. ethylaminothiocarbonylmethylamino, C1-Balkylsulphonylamino, e.g. methylsulphonylamino or ethylsulphonylamino, C1-sdialkylsulphonylamino, e.g. dimethylsulphonyl-amino or diethylsulphonylamino, aminosulphonylamino (-NHSO2NH2), C1-6alkylaminosulphonylamino, e.g. methylaminosulphonylamino or ethylaminosulphonylamino, C1_6dialkylaminosulphonylamino, e.g. dimethylaminosulphonylamino or diethylaminosulphonylamino, C1-salkanoylamino, e.g. acetylamino, aminoC1-salkanoylamino e.g. aminoacetylamino, C1-6dialkylaminoC1-6alkanoylamino, e.g. dimethylaminoacetylamino, C1-6alkanoylaminoC1-6alkyl, e.g. acetylaminomethyl, C1.6alkanoylaminoC1-6alkylamino, e.g. acetamidoethylamino, C1.

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6alkoxycarbonylamino, e.g. methoxycarbonylamino, ethoxycarbonylamino or t-butoxycarbonylamino group.

Optional substituents present on the heteroaromatic groups represented by Het include one, two, three or more substituents, each selected from an atom or group R6 in which R6 is -R6a or -Alk3(R6a), where R6a is a halogen atom, or an amino (-NH2), substituted amino, nitro, cyano, amiding, hydroxyl (-OH) substituted hydroxyl, formyl, carboxyl (-COsH). esterified carboxyl, thiol (-SH), substituted thiol, -CORY (where R7 is an -Alk3(R6a)m, aryl or heteroaryl group], -CSR7, -SO3H, -SO2R7 -SO2NH2, -SO₂NHR⁷ SO₂N(R⁷)₂, -CONH₂, -CSNH₂, -CONHR⁷, -CSNHR⁷. -CONIR7]2, -CSN(R7)2, -N(R4)SO2R7, -N(SO2R7)2, -NH(R4)SO2NH2, -N(R4)SO₂NHR7, -N(R4)SO₂N(R7)₂, -N(R4)COR7, -N(R4)CON(R7)₂, -N(R4)CSN(R7)2, -N(R4)CSR7, -N(R4)C(O)OR7, -SO2NHet1 [where -NHet1 is an optionally substituted Cs.-revelicamino group optionally containing one or more other -O- or -S- atoms or -N(R4)-, -C(O)- or -C(S)- groups], -CONHet1 -CSNHet1. -N(R4)SO₂NHet1. -N(R4)CONHet1, -N(R4)CSNHet1, -SO2N(R4)Het2 [where Het2 is an optionally substituted monocyclic Cs.-zcarbocyclic group optionally containing one or more -O- or -S- atoms or -N(R4)-, -C(O)- or -C(S)- groups], -CON(R4)Het2, -CSN(R4)Het2, -N(R4)CON(R4)Het2,-N(R4)CSN(R4)Het2, anyl or heteroaryl group: Alk3 is a straight or branched C1-salkviene, C2-salkenylene or C2salkynylene chain, optionally interrupted by one, two or three -O- or -Satoms or -S(O)_n [where n is an integer 1 or 2] or -N(R8)- groups [where R8 is a hydrogen atom or C1-salkyl, e.g. methyl or ethyl group); and m is zero or an integer 1, 2 or 3. It will be appreciated that when two R4 or R7 groups are present in one of the above substituents, the R4 or R7 groups may be the same or different.

30 When in the group -Alk³(R^{6a})_m m is an integer 1, 2 or 3, it is to be understood that the substituent or substituents R^{6a} may be present on any suitable carbon atom in -Alk³. Where more than one R^{6a} substituent is present these may be the same or different and may be present on the same or different atom in -Alk³. Clearly, when m is zero and no substituent R^{6a} is present the alkylene, alkenylene or alkynylene chain represented by Alk³ becomes an alkyl, alkenyl or alkynyl group.

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When R^{6a} is a substituted amino group it may be for example a group -NHR⁷ [where R⁷ is as defined above] or a group -N(R⁷)₂ wherein each R⁷ group is the same or different.

When R^{6a} is a halogen atom it may be for example a fluorine, chlorine, bromine, or iodine atom.

When R^{6a} is a substituted hydroxyl or substituted thiol group it may be for 10 example a group -OR⁷ or a -SR⁷ or -SC(=NH)NH₂ group respectively.

Esterified carboxyl groups represented by the group R^{6a} include groups of formula $-CO_2Alk^4$ wherein Alk^4 is a straight or branched, optionally substituted $C_{1-8alkyl}$ group such as a methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl or i-butyl group; a C_{6-12} aryl $C_{1-8alkyl}$ group such as an optionally substituted benzyl, phenylethyl, phenylpropyl, 1-naphthylmethyl or 2-naphthyl group; a C_{6-12} aryl group such as an optionally substituted phenyl, 1-naphthyl or 2-naphthyl group; a C_{6-12} aryloxy $C_{1-8alkyl}$ group such as an optionally substituted phenyloxymethyl, phenyloxymethyl, 1-naphthyl-oxymethyl, or 2-naphthyloxymethyl group; an optionally substituted C_{1-8} alkyl group, such as a pivaloyloxymethyl, propionyloxyethyl or propionyloxypropyl group; or a C_{6-12} aroyloxy C_{1-8} alkyl group such as an optionally substituted benzoyloxyethyl or benzoyloxypropyl group. Optional substituents present on the Alk⁴ group include R^{6a} substituents described above.

When Alk³ is present in or as a substituent it may be for example a methylene, ethylene, n-propylene, i-propylene, n-butylene, i-butylene, s-butylene, t-butylene, ethenylene, 2-propenylene, 2-butynylene, 3-butynylene, 2-propynylene, 2-butynylene or 3-butynylene chain, optionally interrupted by one, two, or three -O- or -S-, atoms or -S(O)-, -S(O)2- or -N(R⁸)- groups.

Aryl or heteroaryl groups represented by the groups R^{6a} or R⁷ include mono- or bicyclic optionally substituted C₆₋₁₂ aromatic or C₁₋₉ heteroaromatic groups as described above for the groups R¹ and Het.

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The aromatic and heteroaromatic groups may be attached to the remainder of the compound of formula (1) by any carbon or hetero e.g. nitrogen atom as appropriate.

5 When -NHet¹ or -Het² forms part of a substituent R⁶ each may be for example an optionally substituted pyrrolidinyl, pyrazolidinyl, piperazinyl, morpholinyl, thiomorpholinyl, piperidinyl or thiazolidinyl group. Additionally Het² may represent for example, an optionally substituted cyclopentyl or cyclohexyl group. Optional substituents which may be present on -NHet¹ or -Het² include those R⁶ substituents described above.

Particularly useful atoms or groups represented by R6 include fluorine. chlorine, bromine or iodine atoms, or C1_salkvl, e.g. methyl, ethyl, n-propyl, i-propyl, n-butyl or t-butyl, optionally substituted phenyl, pyridyl, pyrrolyl, furyl, thiazolyl, or thienyl, C1-6alkylamino, e.g. methylamino or ethylamino, C₁₋₆hydroxyalkyl, e.g. hydroxymethyl or hydroxyethyl, carboxyC₁₋₆alkyl, e.g. carboxyethyl, C1-6alkylthio e.g. methylthio or ethylthio, carboxyC1galkylthio, e.g. carboxymethylthio, 2-carboxyethylthio or 3-carboxypropylthio, C_{1.6}alkoxy, e.g. methoxy or ethoxy, hydroxyC_{1.6}alkoxy, e.g. 2hydroxyethoxy, optionally substituted phenoxy, pyridyloxy, thiazolyloxy, phenylthio or pyridylthio. C5.7cycloalkoxy, e.g. cyclopentyloxy, haloC1. 6alkyl, e.g. trifluoromethyl, haloC1-8alkoxy, e.g. trifluoromethoxy, C1salkylamino, e.g. methylamino or ethylamino, amino (-NH2), aminoC1. salkyl, e.g. aminomethyl or aminoethyl, C1-sdialkylamino, e.g. dimethylamino or diethylamino, C1-6alkylaminoC1-6alkyl, e.g. ethylaminoethyl, C1.edialkylaminoC1.ealkyl, e.g. diethylaminoethyl, aminoC1.ealkoxy, e.g. aminoethoxy, C1-salkylaminoC1-salkoxy, e.g. methylaminoethoxy, C1adialkylaminoC1.ealkoxy, e.g. dimethylaminoethoxy, diethylaminoethoxy, isopropylaminoethoxy, or dimethylaminopropoxy, imido, such as phthalimido or naphthalimido, e.g. 1,8-naphthalimido, nitro, cyano, amidino, hydroxyl (-OH), formyl [HC(O)-], carboxyl (-CO2H), -CO2Alk4 [where Alk4 is as defined above], C1-6 alkanoyl e.g. acetyl, optionally substituted benzoyl, thiol (-SH), thioC1-6alkyl, e.g. thiomethyl or thioethyl, -SC(=NH)NH2, sulphonyl (-SO3H), C1-6alkylsulphoriyl, e.g. methylsulphonyl, optionally substituted phenylsulphonyl, aminosulphonyl (-SO₂NH₂), C₁₋₆alkylaminosulphonyl, e.g. methylaminosulphonyl or

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ethylaminosulphonyl, C1-edialkylaminosulphonyl. dimethylaminosulphonyl or diethylaminosulphonyl, phenylamino-sulphonyl, carboxamido (-CONHa). C1-salkylaminocarbonyl, e.g. methylaminocarbonyl or ethylaminocarbonyl, C1-6dialkylaminocarbonyl, e.g. dimethylaminocarbonyl or diethylaminocarbonyl, aminoC1_6alkylaminocarbonyl, e.g. aminoethylaminocarbonyl, C1-8dialkylaminoC1-8alkylaminocarbonyl, e.g. diethylaminoethylaminocarbonyl, aminocarbonylamino, C₁ salkylaminocarbonylamino, e.g. methylaminocarbonylamino or ethylaminocarbonylamino. C1-sdialkylaminocarbonylamino, e.g. dimethylaminocarbonylamino or diethylaminocarbonylamino. C1-salkylaminocabonylC1salkylamino, e.g. methylaminocarbonylmethylamino, aminothiocarbonylamino. C1.salkvlaminothiocarbonvlamino, e.g. methylaminothiocarbonvlamino or ethylaminothiocarbonylamino, C1-8dialkylaminothiocarbonylamino, e.g. dimethylaminothiocarbonylamino or diethylaminothiocarbonylamino. C1-salkylaminothiocarbonylC1-salkylamino, e.g. ethylaminothiocarbonylmethylamino, -CONHC(=NH)NH2, C1-6alkylsulphonylamino, e.g. methylsulphonylamino or ethylsulphonylamino, C1-sdialkylsulphonylamino, e.g. dimethylsulphonylamino or diethylsulphonylamino, optionally substituted phenylsulphonylaming, aminosulphonylaming (-NHSO2NH2), C1-6alkylaminosulphonylamino, e.g. methylaminosulphonyl-amino or ethylaminosulphonylamino, C1-6dialkylaminosulphonylamino, e.g. dimethylaminosulphonylamino or diethylaminosulphonylamino, optionally substituted morpholinesulphonylamino or morpholinesulphonylC1-salkylamino, optionally substituted phenylaminosulphonylamino, C1.ealkanoviamino, e.g. acetylamino, aminoC1-6alkanoylamino e.g. aminoacetylamino, C1_sdialkylaminoC1_salkanovl-amino, e.g. dimethylaminoacetylamino, C1-6alkanoylaminoC1-6alkyl, e.g. acetylaminomethyl, C1-6alkanoylaminoC1salkylamino, e.g. acetamidoethylamino, C1-5alkoxycarbonylamino, e.g. methoxycarbonylamino, ethoxycarbonylamino or t-butoxycarbonylamino or optionally substituted benzyloxy, pyridylmethoxy, thiazolylmethoxy, benzyloxycarbonylamino, benzyloxycarbonylaminoC1-6alkyl e.g. benzyloxycarbonylaminoethyl, benzylthio, pyridylmethylthio or thaizolylmethylthio groups.

In the above groups of particularly useful R⁶ substituents, the reference to optional substitution is intended to relate primarily to the aromatic or

heteroaromatic portions of the groups described. Thus for example such groups may be optionally mono-, di- or tri-substituted by those particular atoms or groups described above for each of R^a and R^b.

5 Where desired, two R⁶ substituents may be linked together to form a cyclic group such as a cyclic ether, e.g. a C_{1.6}alkylenedioxy group such as methylenedioxy or ethylenedioxy.

It will be appreciated that where two or more R6 substituents are present, these need not necessarily be the same atoms and/or groups. In general, the substituent(s) may be present at any available ring position in the heteroaromatic group represented by Het.

The presence of certain substituents in the compounds of formula (1) may enable salts of the compounds to be formed. Suitable salts include pharmaceutically acceptable salts, for example acid addition salts derived from inorganic or organic acids, and salts derived from inorganic and organic bases.

20 Acid addition salts include hydrochlorides, hydrobromides, hydroiodides, alkylsulphonates, e.g. methanesulphonates, ethanesulphonates, or isothionates, arylsulphonates, e.g. p-toluenesulphonates, besylates or napsylates, phosphates, sulphates, hydrogen sulphates, acetates, trifluoroacetates, propionates, citrates, maleates, fumarates, malonates, succinates, lactates, oxalates, tartrates and benzoates.

Salts derived from inorganic or organic bases include alkali metal salts such as sodium or potassium salts, alkaline earth metal salts such as magnesium or calcium salts, and organic amine salts such as morpholine, piperidine, dimethylamine or diethylamine salts.

Particularly useful saits of compounds according to the invention include pharmaceutically acceptable saits, especially acid addition pharmaceutically acceptable saits.

R in compounds of the invention is preferably a -CO₂H group.

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When present, the aliphatic chain represented by Alk¹ in compounds of the invention is preferably a -CHo- chain.

- 5 Alk² in compounds of formula (1) is preferably a -CH₂- chain and m is preferably an integer 1. In compounds of this type, the carbon atom to which Alk² and R are attached forms a chiral centre and is preferably in the L configuration.
- 10 R² in compounds of formula (1) is preferably a hydrogen atom.

R3 in compounds of the invention is preferably a hydrogen atom.

In general in compounds of the invention - $(Alk^1)_s$ is preferably - CH_2O -,- SO_2NH -, -C(O)O- or - $CON(R^4)$ - and is especially -CONH-.

In general in compounds of the invention the group R^1 is preferably an optionally substituted aromatic or heteroaromatic group. Particularly useful groups of these types include optionally substitued phenyl, pyridyl or pyrimidinyl groups, particularly those in which the substituent when present is an atom or group $-L^2(CH_2)_pL^3(R^0)_q$ as described above. Each substituent may be present on any available ring carbon or nitrogen atom.

The heteroaromatic group represented by Het in compounds of formula (1) is preferably on optionally substituted C_{3-5} monocyclic heteroaromatic group containing one, two or three heteroatoms selected from oxygen, sulphur or nitrogen atoms. Particularly useful groups of this type include optionally substituted pyrrolyl and pyridyl groups. Especially useful heteroaromatic groups represented by Het include optionally substituted 3-or 4-pyridyl groups, particularly 2-monosubstituted 3- or 4-pyridyl or 2.6-disubstituted 3- or 4-pyridyl groups. In these, and in general in the group Het, the optional substituent when present is preferably an atom or group R^6 as defined above.

35 A particularly useful class of compounds according to the invention has the formula (1a)

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$$\begin{array}{c} \mathbb{R}^9 \\ \text{VAIK}^1)_{\text{r}}(\mathbb{L}^1)_{\text{e}} & \mathbb{R}^4 \\ \mathbb{R}^{10} & \mathbb{R}^5 & \mathbb{C}H_2 \\ \mathbb{R} & \mathbb{R}^5 \end{array}$$

wherein -W= is -CH= or -N=, R9 and R10, which may be the same or different is each a -L2(CH2)0L3(Rc)0 atom or group as generally and particularly defined above, and Alk1, r, L1, s, Ra, Rb, R and Het are as generally and particularly defined above, and the salts, solvates, hydrates and N-oxides thereof.

10 It will be appreciated that the various preferences stated above in relation to groups present in compounds of formula (1) apply equally to the same groups when present in compounds of formula (1).

Additionally, in the compounds of formula (1a) -(Alk1)r(L1)s- is preferably a -CH2O or -CON(R4)- group and is especially a -CONH- group. Het is preferably an optionally substituted pyrrolyl or especially an optionally substitued pyridyl group.

Particularly useful compounds of formula (1a) are those wherein Het is a 2-monosubstituted 3- or 4-pyridyl group or a 2,6-disubstituted 3- or 4pyridyl group.

One of R9 or R10 in compounds of formula (1a) may be for example a hydrogen atom and the other a substituent L2(CH2)pL3(Rc)g in which Rc is not a covalent bond and p is zero, but preferably each of R9 and R10 is a substituent -L(2CH2)pL3(Rc)q where Rc is as just defined. Particularly useful R9 or R10 substituents include a hydrogen atom or halogen atom. especially fluorine or chlorine atoms, or a methyl, ethyl, methoxy, ethoxy, -CF₃, -OH, -CN, -NO₂, -NH₂, -NHCH₂, -N(CH₃)₂, -COCH₃, -SCH₃, -CO₂H or -CO₂CH₃ group.

Particularly useful compounds according to the invention include the following:

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- 2-Thio(S-2,5-dimethoxyphenyl)nicotinoyl-(*N*-2,6-dichlorobenzoyl)-*L*-4-aminophenylalanine;
- 2-Thio(S-2,5-dimethoxyphenyl)nicotinoyl-(*N*-2,6-dichlorobenzoyl)-*L*-4-aminophenylalanine;
- 5 N-(3,5-Dichloro-4-picolyl)-N-(3,5-dichloro-4-picolyl)-L-4-aminophenylalanine:
 - N-(2-Chloronicotinoyl)-N-(3,5-dichloro-4-picolyl)-L-4-amino-phenylalanine; O-(2,6-dichlorobenzyl)-N-(4-acetyl-1,2,5-trimethyl-3-pyrroyl)-L-tyrosine;
 - (*N*'-3,5-Dichloroisonicotinoyl)-*N*-{([3-pyridinylmethyl]thio)isonicotinoyl}-*L*-4-aminophenylalanine:
 - N-(4-Acetyl-1,2,5-trimethyl-1/+pyrrole-3-carbonyl)-N-(3,5-dichloro-4picolyl)-L-4-aminophenylalanine; and the salts, solvates: hydrates and Noxides thereof.
- 15 Compounds according to the invention are potent and selective inhibitors of α4 integrins. The ability of the compounds to act in this way may be simply determined by employing tests such as those described in the Examples hereinafter. In particular compounds of the invention, such as the compounds of formula (1a) herein, the compounds are advantageously selective α481 inhibitors.
 - The compounds are of use in modulating cell adhesion and in particular are of use in the prophylaxis and treatment of diseases or disorders involving inflammation in which the extravasation of leukocytes plays a role and the invention extends to such a use and to the use of the compounds for the manufacture of a medicament for treating such diseases or disorders.
- Diseases or disorders of this type include inflammatory arthritis such as 30 rheumatoid arthritis vasculitis or polydermatomyositis, multiple sclerosis, allograft rejection, diabetes, inflammatory dermatoses such as psoriasis or dermatitis, asthma and inflammatory bowel disease.
 - For the prophylaxis or treatment of disease the compounds according to the invention may be administered as pharmaceutical compositions, and according to a further aspect of the invention we provide a pharmaceutical

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composition which comprises a compound of formula (1) together with one or more pharmaceutically acceptable carriers, excipients or diluents.

Pharmaceutical compositions according to the invention may take a form suitable for oral, buccal, parenteral, nasal, topical or rectal administration, or a form suitable for administration by inhalation or insufflation.

For oral administration, the pharmaceutical compositions may take the form of, for example, tablets, lozenges or capsules prepared by conventional means with pharmaceutically acceptable excipients such as binding agents (e.g. pregelatinised maize starch, polyvinylpyrrolidone or hydroxypropyl methylcellulose); fillers (e.g. lactose, microcrystalline cellulose or calcium hydrogen phosphate); lubricants (e.g. magnesium stearate, talc or silica); disintegrants (e.g. potato starch or sodium glycollate); or wetting agents (e.g. sodium lauryl sulphate). The tablets may be coated by methods well known in the art. Liquid preparations for oral administration may take the form of, for example, solutions, syrups or suspensions, or they may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations may be prepared by conventional means with pharmaceutically acceptable additives such as suspending agents, emulsifying agents, non-aqueous vehicles and preservatives. The preparations may also contain buffer salts, flavouring, colouring and sweetening agents as appropriate.

25 Preparations for oral administration may be suitably formulated to give controlled release of the active compound.

For buccal administration the compositions may take the form of tablets or lozences formulated in conventional manner.

The compounds for formula (1) may be formulated for parenteral administration by injection e.g. by bolus injection or infusion. Formulations for injection may be presented in unit dosage form, e.g. in glass ampoule or multi dose containers, e.g. glass vials. The compositions for injection may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as

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suspending, stabilising, preserving and/or dispersing agents. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, e.g., sterile pyrogen-free water, before use.

- 5 In addition to the formulations described above, the compounds of formula (1) may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation or by intramuscular injection.
- For nasal administration or administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation for pressurised packs or a nebuliser, with the use of suitable propellant, e.g. dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dloxide or other suitable use or mixture of cases.

The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient. The pack or dispensing device may be accompanied by instructions for administration.

The quantity of a compound of the invention required for the prophylaxis or treatment of a particular condition will vary depending on the compound chosen, and the condition of the patient to be treated. In general, however, daily dosages may range from around 100ng/kg to 100mg/kg e.g. around 0.01mg/kg to 40mg/kg body weight for oral or buccal administration, from around 10ng/kg to 50mg/kg body weight for parenteral administration and around 0.05mg to around 1000mg e.g. around 0.5mg to around 1000mg for nasal administration or administration by inhalation or insufflation.

The compounds of the invention may be prepared by a number of processes as generally described below and more specifically in the Examples hereinafter. In the following process describtion, the symbols R, R1-R3, Ra, Rb, L1, Alk1, Alk2, m, r, s and Het when used in the formulae depicted are to be understood to represent those groups described above

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in relation to formula (1) unless otherwise indicated. In the reactions described below, it may be necessary to protect reactive functional groups, for example hydroxy, amino, thio or carboxy groups, where these are desired in the final product, to avoid their unwanted participation in the reactions. Conventional protecting groups may be used in accordance with standard practice [see, for example, Green, T. W. in "Protective Groups in Organic Synthesis", John Wiley and Sons, 1991]. In some instances, deprotection may be the final step in the synthesis of a compound of formula (1) and the processes according to the invention described hereinafter are to be understood to extend to such removal of protecting groups.

Thus according to a further aspect of the invention, a compound of formula (1) may be obtained by hydrolysis of an ester of formula (2):

$$R^{1}(A|k^{1})_{i}(L^{1})_{5}$$
 R^{0}
 $(A|k^{2})_{m}$
 $C(R^{2})$
 $CO_{2}R^{11}$
(2)

where R11 is an alkyl group.

The hydrolysis may be performed using either an acid or a base depending on the nature of R¹¹, for example an organic acid such as trifluoracetic acid or an inorganic base such as lithium hydroxide optionally in an aqueous organic solvent such as an amide, e.g. a substituted amide such as dimethylformamide, an ether, e.g. a cyclic ether such as tetrahydrofuran or dioxane or an alcohol, e.g. methanol at around ambient temperature. Where desired, mixtures of such solvents may be used.

Esters of formula (2) may be prepared by coupling an amine of formula (3):

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(where R11 is as just described) or a salt thereof with an acid of formula (4):

HetCO₂H (4)

or an active derivative thereof.

10 Active derivatives of acids of formula (4) include anhydrides, esters and halides. Particular esters include pentafluorophenyl or succinyl esters.

The coupling reaction may be performed using standard conditions for reactions of this type. Thus for example the reaction may be carried out in a solvent, for example an inert organic solvent such as an amide, e.g. a substituted amide such as dimethylformamide, an ether, e.g. a cyclic ether such as tetrahydrofuran, or a halogenated hydrocarbon, such as dichloromethane, at a low temperature, e.g. around -30°C to around ambient temperature, optionally in the presence of a base, e.g. an organic base such as an amine, e.g. triethylamine, pyridine, or dimethylaminopyridine, or a cyclic amine, such as N-methylmorpholine.

Where an acid of formula (4) is used, the reaction may additionally be performed in the presence of a condensing agent, for example a diimide such as 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide or N,N'-dicyclohexylcarbodiimide, advantageously in the presence of a catalyst such as a N-hydroxy compound e.g. a N-hydroxytriazole such as 1-hydroxybenzotriazole. Alternatively, the acid may be reacted with a chloroformate, for example ethylchloroformate, prior to reaction with the amine of formula (3).

Intermediates of formulae (2), (3) and (4), or compounds of formula (1), may be manipulated to introduce substituents to aromatic or heteroaromatic groups or modify existing substituents in groups of these types. Typically, such manipulation may involve standard substitution approaches employing for example alkylation, arylation, heteroarylation, acylation, thioacylation, halogenation, suphonylation, nitration, formylation or coupling reactions. Alternatively, existing substituents may be modified for example by oxidation, reduction or cleavage reactions. Particular examples of such reactions are given below. Where these are described in relation to the generation of the group R¹(Alk¹),(L¹)s⁻, it will be appreciated that each reaction may also be used to introduce or modify R⁵ and/or R⁶ substituents as appropriate.

Thus in one example, a compound wherein $H^1(Alk^1)_r(L^1)_{S^*}$ is a -L¹H group may be alkylated, arylated or heteroarylated using a reagent $H^1(Alk^1)_rX$ in which H^1 is other than a hydrogen atom and X is a leaving atom or group such as a halogen atom, e.g. a fluorine, bromine, iodine or chlorine atom or a sulphonyloxy group such as an alkylsulphonyloxy, e.g. trifluoromethylsulphonyloxy or arylsulphonyloxy, e.g. p-toluenesulphonyloxy group.

The reaction may be carried out in the presence of a base such as a carbonate, e.g. caesium or potassium carbonate, an alkoxide, e.g. potassium t-butoxide, or a hydride, e.g. sodium hydride, in a dipolar aprotic solvent such as an amide, e.g. a substituted amide such as dimethylformamide or an ether, e.g. a cyclic ether such as tetrahydrofuran.

In another example, a compound where R¹(Alk¹),(L¹)₈ is a -L¹H group is a hydrogen atom may be functionalised by acylation or thioacylation, for example by reaction with a reagent R¹(Alk¹),L¹X [wherein L¹ is a -C(O)-, C(S)-, -N(R⁴)C(O)- or N(R⁴)C(S)- group], in the presence of a base, such as a hydride, e.g. sodium hydride or an amine, e.g. triethylamine or N-methylmorpholine, in a solvent such as a halogenated hydrocarbon, e.g. dichloromethane or carbon tetrachloride or an amide, e.g. dimethylformamide, at for example ambient temperature, or by reaction

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with R¹(Alk¹),CO₂H, R¹(Alk)₄COSH or an activated derivative thereof, for example as described above for the preparation of esters of formula (2).

In a further example a compound may be obtained by sulphonylation of a compound where $R^1(Alk^1)_f(L^1)_s$ is an -OH group by reaction with a reagent $R^1(Alk^1)_fL^1Hal$ [in which L^1 is -S(O)- or -SO2- and Hal is a halogen atom such as chlorine atom] in the presence of a base, for example an inorganic base such as sodium hydride in a solvent such as an amide, e.g. a substituted amide such as dimethylformamide at for example ambient temperature.

In another example, a compound where $R^1(Alk^1)_A(L^1)_8$ is a -L¹H group, may be coupled with a reagent R^1OH (where R^1 is other than a hydrogen atom) or R^1Alk^1OH in a solvent such as tetrahydrofuran in the presence of a phosphine, e.g. triphenylphosphine and an activator such as diethyl, diisopropyl- or dimethylazodicarboxylate to yield a compound containing a $R^1(Alk^1)_O$ - group.

In a further example, ester groups -CO₂R4 or -CO₂Alk4 in compounds of formula (1) may be converted to the corresponding acid [-CO₂H] by acid-or base-catalysed hydrolysis depending on the nature of the grousp R4 or Alk4. Acid- or base-catalysed hydrolysis may be achieved for example by treatment with an organic or inorganic acid, e.g. trifluoroacetic acid in an aqueous solvent or a mineral acid such as hydrochloric acid in a solvent such as dioxan or an alkali metal hydroxide, e.g. lithium hydroxide in an aqueous alcohol, e.g. aqueous methanol.

In a second example, -OR⁷ [where R⁷ represents an alkyl group such as methyl group] groups in compounds of formula (1) may be cleaved to the corresponding alcohol -OH by reaction with boron tribromide in a solvent such as a halogenated hydrocarbon, e.g. dichloromethane at a low temperature, e.g. around -78°C.

Alcohol [-OH] groups may also be obtained by hydrogenation of a corresponding -OCH₂R⁷ group (where R⁷ is an aryl group) using a metal catalyst, for example palladium on a support such as carbon in a solvent

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such as ethanol in the presence of ammonium formate, cyclohexadiene or hydrogen, from around ambient to the reflux temperature. In another example, -OH groups may be generated from the corresponding ester [-CO₂Alk⁴ or CO₂Alk⁴] or aldehyde [-CHO] by reduction, using for example a complex metal hydride such as lithium aluminium hydride or sodium borohydride in a solvent such as methanol.

In another example, alcohol -OH groups in compounds of formula (1) may be converted to a corresponding -OR³ group by coupling with a reagent R⁷OH in a solvent such as tetrahydrofuran in the presence of a phosphine, e.g. triphenylphosphine and an activator such as diethyl-, diisopropyl-, or dimethylazodicarboxvlate.

Aminosulphonylamino [-NHSO₂NH₂] groups in compounds of formula (1) may be obtained, in another example, by reaction of a corresponding amine [-NH₂] with sulphamide in the presence of an organic base such as pyridine at an elevated temperature, e.g. the reflux temperature.

In a further example amine (-NH₂) groups may be alkylated using a reductive alkylation process employing an aldehyde and a borohydride, for example sodium triacetoxyborohyride or sodium cyanoborohydride, in a solvent such as a halogenated hydrocarbon, e.g. dichloromethane, a ketone such as acetone, or an alcohol, e.g. ethanol, where necessary in the presence of an acid such as acetic acid at around ambient temperature.

In a further example, amine [-NH₂] groups in compounds of formula (1) may be obtained by hydrolysis from a corresponding imide by reaction with hydrazine in a solvent such as an alcohol, e.g. ethanol at ambient temperature.

In another example, a nitro [-NO₂] group may be reduced to an amine [-NH₂], for example by catalytic hydrogenation using for example hydrogen in the presence of a metal catalyst, for example palladium on a support such as carbon in a solvent such as an ether, e.g. tetrahydrofuran or an

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alcohol e.g. methanol, or by chemical reduction using for example a metal, e.g. tin or iron, in the presence of an acid such as hydrochloric acid.

Aromatic halogen substituents in compounds of the invention may be subjected to halogen-metal exchange with a base, for example a lithium base such as n-butyl or t-butyl lithium, optionally at a low temperature, e.g. around -78°C, in a solvent such as tetrahydrofuran and then quenched with an electrophile to introduce a desired substituent. Thus, for example, a formyl group may be introduced by using dimethylformamide as the electrophile; a thiomethyl group may be introduced by using dimethyldisulohide as the electrophile.

In another example, sulphur atoms in compounds of the invention, for example when present in the linker group L¹ may be oxidised to the corresponding sulphoxide using an oxidising agent such as a peroxy acid, e.g. 3-chloroperoxybenzoic acid, in an inert solvent such as a halogenated hydrocarbon, e.g. dichloromethane, at around ambient temperature.

Intermediates of formulae (3) and (4), R¹(Alk¹)_rX, R¹(Alk¹)_rL¹X, R¹(Alk¹)_rCO₂H, R¹OH and R¹Alk¹OH are either known compounds or may be prepared from known starting materials by use of analogous processes to those used for the preparation of the known compounds and/or by treating known compounds by one or more of the alkylation, acylation and other manipulations described herein, such as particularly described for the preparation of the Intermediates in the exemplification selection hereinafter.

N-oxides of compounds of formula (1) may be prepared for example by oxidation of the corresponding nitrogen base using an oxidising agent such as hydrogen peroxide in the presence of an acid such as acetic acid, at an elevated temperature, for example around 70°C to 80°C, or alternatively by reaction with a peracid such as peracetic acid in a solvent, e.g. dichloromethane, at ambient temperature.

35 Salts of compounds of formula (1) may be prepared by reaction of a compound of formula (1) with an appropriate base in a suit able solvent or

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mixture of solvents e.g. an organic solvent such as an ether e.g. diethylether, or an alcohol, e.g. ethanol using conventional procedures.

Where it is desired to obtain a particular enantiomer of a compound of 5 formula (1) this may be produced from a corresponding mixture of enantiomers using any suitable conventional procedure for resolving enantiomers.

Thus for example diastereomeric derivatives, e.g. salts, may be produced by reaction of a mixture of enantiomers of formula (1) e.g. a racemate, and an appropriate chiral compound, e.g. a chiral base. The diastereomers may then be separated by any convenient means, for example by crystallisation and the desired enantiomer recovered, e.g. by treatment with an acid in the instance where the diastereomer is a salt.

In another resolution process a racemate of formula (1) may be separated using chiral High Performance Liquid Chromatography. Alternatively, if desired a particular enantiomer may be obtained by using an appropriate chiral intermediate in one of the processes described above.

The following Examples illustrate the invention. All temperatures are in °C. The following abbreviations are used:

EDC - 1-(3-dimethylaminopropyl)3-ethycarbodiimide:

DMF - dimethylformamide;

HOBT - 1-hydroxybenzotriazole; TFA - trifluoroacetic acid:

DCM - dichloromethane:

BOC - tert-butoxycarbonyl;
30 MeOH - methanol;

tvr - tvrosine:

HetAr - heteroaryl; thiopro - thioproline;

Me - methyl:

DMSO - dimethylsulphoxide;

THF - tetrahydrofuran;

NMM - N-methylmorpholine;

Ph - phenyl;

EtOAc - ethyl acetate;

LDA - lithium diisopropylamide

Ar - aryl;

pyr - pyridine;

Bu - butyl;

app - apparent

INTERMEDIATE 1

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2-Chloronicotinovi-O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester

A solution of O-(2,6-dichlorobenzyl)-*L*-tyrosine methyl ester hydrochloride, (1.11g, 2.84mmol), 2-chloronicotinic acid (0.45g, 2.84mmol), EDC (0.60g, 3.13mmol), HOBT (0.46g, 3.41mmol) and NMM (0.467ml, 0.43g, 4.26mmol) in DCM (25ml) was stirred at room temperature for 24 h. The reaction mixture was partitioned between DCM (50ml) and 10% NaHCO₃ solution (30ml). The organic layer was separated, dried over MgSO₄ and the solvent removed under vacuum to give a pale yellow solid that was recrystalised from EtOAc/hexane to give the title compound as an off white solid (1.14g, 81%). 8H (CDCl₃) 8.46 (1H, dd, ½ 2.0, 4.7Hz), 8.05 (1H, dd, ½ 2.0, 7.6Hz), 7.34 (2H, m), 7.25 (2H, m), 7.11 (2H, m), 6.96 (3H, m), 5.24 (2H, s), 5.06 (1H, m), 3.80 (3H, s) and 3.24 (2H, m), 6.96 (3H, m), 5.24

15 INTERMEDIATE 2

2-Thio(\$-2.5-dimethoxyphenyl)nicotinoyi-Q-(2,6-dichlorobenzyl)-L-tyrosine methyl ester

A solution of O-(2,6-dichiorobenzyl)-L-tyrosine methyl ester hydrochloride (0.78g, 2.0mmol), EDC (0.42g, 2.2mmol) HOBT (0.32g, 2.4mmol) and NMM (0.55ml, 0.50g, 5.0mmol) in DMF (10ml) was treated with a solution of 2-thio(2,5-dimethoxyphenyl)nicotinic acid (0.58g, 2.0mmol) in DMF (2ml) and stirred for 16h at room temperature. Solvent was removed in vacuo and the residue was partitioned between EtOAc (55ml) and 10% hydrochloric acid (25ml). The organic layer was separated, washed with 10% NaHCO₃ solution (30ml), dried over MgSO₄ and the solvent removed in vacuo to give a yellow oil which was purified by chromatography (SiO₂: EtOAc/hexane 1:1) to give the title compound as a white foam (1.14g, 86%). SH (CDCi₃) 8.39 (1H, dd, ½ 1.9, 4.7Hz), 7.85 (1H, dd, ½ 1.9, 7.7Hz), 7.47 (1H, d, ½ 7.6HJz), 7.37-6.81 (11H, m), 5.18 (2H, s), 5.09 (1H, m), 3.78 (6H, s), 3.53 (3H, s) and 3.25 (2H, m).

INTERMEDIATE 3

2-Mercaptonicotinoyi-*O*-(2,6-dichlorobenzyi)-*L*-tyrosine methyl ester

A solution of O-(2,6-dichlorobenzyi)-*L*-tyrosine methyl ester hydrochloride
(2.50g, 6.4mmol), 2-mercaptonicotinic acid (0.99g, 6.4mmol) and NMM
(1.41ml, 1.29g, 12.8mmol) in DMF (10ml) was stirred at room temperature

for 64h. Solvent was removed *in vacuo* and the residue partitioned between DCM (30ml) and water (25ml). The aqueous layer was extracted with DCM (30ml) and the combined organic layers were washed with 10% NaHCO₃ solution (30ml), dried over MgSO₄ and the solvent removed *in vacuo* to give a brown oil which was purified by chromatography (SiO₂; gradient elution, 4:1 EtOAc/hexane to 100% EtOAc) to give the title compound as a yellow foam, (2.92g, 93%). 8H (CDCl₃) 8.71 (1H, dd, J. 1.8, 7.6Hz), 8.05 (1H, s), 7.61 (1H, dd, J. 1.8, 6.1Hz), 7.35 (2H, m), 7.33-7.19 (2H, m), 6.94 (2H, d, J. 8.7Hz), 5.22 (2H, s), 4.97 (1H, m), 3.74 (3H, s) and 3.21 (2H, m).

INTERMEDIATE 4

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2-Thio(S-4-picolinyl)nicotinoyl-O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester

A solution of Intermediate 3 (0.50g, 1.0mmol) and 4-picolyl chloride hyrochloride (0.17g, 1.0mmol) in DCM (10ml) was treated with 1,8-diazabicyclo[5.4.0]undec-7-ene (0.31ml, 0.31g, 2.0mmol) and stirred at room temperature for 5h. The reaction was partitioned between water and DCM, the organic layer separated, dried over MgSO₄ and the solvent removed *in vacuo* to give a yellow gum that was purified by chromatography (SiO₂, EtOAc), to give a pale yellow solid, which was recrystallised from EtOAc/nexane (1:1) to give the <u>title compound</u> as an off white solid (0.30g, 52%). 5H (CDCl₃) 8.47(2H, m), 7.69 (1H, dd, <u>J</u> 1.8, 77Hz), 7.38-7.22 (6H, m), 7.06 (3H, m), 6.90 (2H, m), 6.62 (1H, d, <u>J</u> 7.5Hz), 5.23 (2H, s), 5.03 (1H, m), 4.40 (2H, m), 3.79 (3H, s), 3.28 (1H, dd, <u>J</u> 5.8, 14.1Hz) and 3.19 (1H, dd, <u>J</u> 5.4, 14.1Hz).

INTERMEDIATE 5

2-Thio(S-2.5-dimethoxyphenyl)nicotinoyl-L-4-aminophenylalanine methyl ester

A solution of 4-amino-L-phenylalanine methyl ester dihydrochloride (0.53g, 2.0mmol), EDC (0.42g, 2.2mmol) HOBT (0.32g, 2.4mmol) and NMM (0.66ml, 0.61g, 6.0mmol) in DMF (10ml) was treated with a solution of 2-thio(2,5-dimethoxyphenyl)nicotinic acid (0.58g, 2.0mmol) in DMF (2ml) and stirred for 64h at room temperature. The solvent was removed in vacuo, and the residue partitioned between DCM (30ml) and water (20ml). The

organic layer was separated, washed with 10% NaHCO₃ (20ml) solution, dried over MgSO₄ and the solvent evaporated *in vacuo* to give a brown gum which was purified by chromatography (SiO₂; EtOAc) to give the <u>title compound</u> as a yellow foam (0.67g, 72%). 8H (CDO₃) 8.35 (1H, dd, <u>1</u> 1.8, 4.8Hz), 7.81 (1H, dd, <u>1</u> 1.9, 7.7Hz), 7.41 (1H, d, <u>1</u> 7.6Hz), 7.12 (1H, d, <u>1</u> 3.0Hz), 7.07-6.81 (5H, m), 6.49)2H, d, <u>1</u> 8.4Hz), 5.00 (1H, m), 3.76 (3H, s), 3.74 (3H, s, 3.55 (3H, s) and 3.14 (2H, m).

INTERMEDIATE 6

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10 2-Thio(\$-2.5-dimethoxyphenyl)nicotinoyl-(N-2.6-dichlorobenzoyl)-L-4aminophenylatanine methyl ester

A solution of Intermediate 5 (0.68g, 1.5mmol) and NMM (0.53ml, 0.49g, 4.8mmol) in DCM (20ml) was treated with 2,6-dichlorobenzoyl chloride (0.23ml, 0.33g, 1.6mmol) and the reaction stirred for 16h at room temperature, then partitioned between DCM (50ml) and 10% NaHCO₃ solution (30ml). The organic layer was separated, dried over MgSO₄ and the solvent evaporated *in vacuo* to give an off-white solid that was triturated with EtOAc/dlethyl ether (2:1) to give the <u>title compound</u> as an off-white solid (0.44g, 46%). δH (MeOH-d⁴) 8.28 (1H, dd, <u>J</u> 1.8, 4.9Hz), 7.70 (1H, dd, <u>J</u> 1.8, 7.6Hz), 7.58 (2H, d, <u>J</u> 8.6Hz), 7.48-7.38 (3H, m), 7.30 (2H, d, <u>J</u> 8.6Hz), 7.27 (1H, dd, <u>J</u> 4.9, 7.7Hz), 7.01 (1H, dd, <u>J</u> 1.1, 2.3Hz), 6.93 (2H, m), 4.87 (1H, m), 3.77 (3H, s), 3.75 (3H, s), 3.80 (3H, s), 3.32 (1H, m) and 3.13 (H, dd, J 8.6, 14.0Hz).

25 INTERMEDIATE 7

<u>O-(2,6-dichlorobenzyl)-N-(4-acetyl-1,2,5-trimethyl-3-pyrroyl)-L-tyrosine methyl ester</u>

NMM (155mg, 169µl, 1.54mmol), HOBT (227mg, 1.58mmol), 4-acetyl-1,2,5-trimethylpyrrole-3-carboxylic acid (300mg, 1.54mmol) and EDC (295mg, 1.54mmol) were added sequentially to a stirred solution of *O*-(2,6-dichlorobenzyl)-*L*-tyrosine methyl ester hydrochloride (546mg, 1.40mmol) in dry DMF (15ml). The reaction was stirred at room temprature under N₂ for 18h. The solvent was removed *in vacuo* and the residue partitioned between EtOAc (50ml), and 10% aqueous Na₂CO₃ (40ml). The phases were separated and the queous phase extracted with EtOAc (2 x 25ml). The combined organic extracts were washed consecutively with 5%

aqueous hydrochloric acid (20ml), 10% aqueous Na₂CO₃ (20ml) and brine (10ml), dried (Na₂SO₄) and evaporated *in vacuo*. The obtained orange foam (0.6g) was chromatographed (silica; 50% EtOAc/Hexane —> 100% EtOAc) affording the <u>title compound</u> as a white foam (380mg, 51%); 'Hnmr (d⁶DMSO) 8.53 (1H, d, \downarrow 8Hz, NH), 7.57-7.43 (3H, m's, aryH-H), 7.22 (2H, d, \downarrow 8.5Hz), aryl-H), 6.95 (2H, d, \downarrow 8.5Hz, aryl-H), 5.18 (2H, br s, CH₂-O), 4.67 (1H, m, α -tyr-H), 3.66 (3H, s, Me-O), 3.34 (3H, s, Me-N), 3.12 (1H, dd, \downarrow 4.1, 13.8Hz, CH_AHgAr), 2.90 (1H, dd, \downarrow 11.3, 13.8Hz, CH_AHgAr), 2.92 (3H, s, Me-O), 2.0 (3H, s, pyrrole-Me) and 1.94 (3H, s, pyrrole-Me). m/z (ES + 60V) 531 (MH+, 100), 533 (MH+, 75) 553 (MNa+, 15%).

INTERMEDIATE 8

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2-Chloronicotinyl-(*N*-2,6-dichlorobenzoyl)-*L*-4-aminophenylalanine methyl ester

EDC (270mg, 1.5mmol) was added to a stirred solution of (N-2,6-dichlorobenzoyl)-L-4-aminophenylalanine methyl ester (500mg, 1.3mmol), 2 chloronicotinic acid (200mg, 1.3mmol), HOBT (190mg, 1.5mmol) and NMM (423µl, 3.9mmol) in anhydrous DMF (2ml) at 0°. The DMF solution was stirred overnight at room temeprature then the DMF was evaporated in vacuo. The residue was taken up in DCM (50ml), washed with water (3 x 10ml), saturated aqueous NaHCO₃ (2 x 10ml) and water (2 x 10ml), dried (Na₂SO₄) and evaporated in vacuo. The residue was purified by chromatography (SiO₂; 1:1 EtOAc: hexane) to give the title compound as a white foam (450mg, 80%). 3H (CDCl₃), 8.50 (1H, m, pyr H), 8.05 (1H, d, ½ 9.2Hz, pyrH), 7.58 (2H, d, ½ 8.5Hz, ArH), 7.48 (1H, br s, NH), 7.36-7.24 (3H, m, 2ArH, 1pyrH), 7.21 (2H, d, ½ 8.5Hz, 2 ArH), 7.12 (1H, d, ½ 6.6Hz, M.), 5.05-5.18 (1H, m, CHottyr), 3.81 (3H, s, CO₂Me) and 3.39-3.18 (2H, m, CH₂Ar).

30 INTERMEDIATE 9

Methyl-2-thio(S-acetate)nicotinoyi-Q-(2,6-dichlorobenzyl)-L-tyrosine methyl ester

A solution of Intermediate 3 (370mg, 0.76mmol) in anhydrous DMF (2ml) was added to a suspension of sodium hydride (60% in oil, 33mg, 0.83mmol) in anhydrous DMF (3ml) at 0°. The mixture was stirred for 10min at room temperature, then recooled to 0°. Methyl bromoacetate

(115mg, 0.76mmol) was added dropwise, then the mixture was stirred overnight at room temperature. The mixture was guenched with water (0.5ml) and the DMF evaporated in vacuo. The residue was dissolved in EtOAc, washed with water (3 x 10ml), dried (Na>SO₄) and evaporated in vacuo. The residue was purified by chromatography (SiO₂: EtOAc/hexane 1:1) to give the title compound as a white solid (360mg, 85%). δH (CDCl₃) 8.45 (1H, m. pvrH), 7.71 (1H, d, J 7.7, pvrH), 7.35 (2H, d, J 7.3Hz, 2ArH), 7.28-6.9 (6H, m, 5ArH, 1pyrH), 6.71 (1H, d, J 7.5Hz, NH), 5.24 (2H, s, OCH2Ar), 5.08-5.03 (1H, m, CHa tvr), 3.96 (2H, s, SCH2), 3.78 (3H, s, CO₂CH₃), 3.71 (3H, s, CO₂CH₃), 3.31 (1H, dd, J 14, 5.4Hz, CH_AH_BAr) and 3.21 (1H, dd, J 14, 5,2Hz, CHAHRAr).

INTERMEDIATE 10

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2-Thio(S-methyl)nicotinovi-O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester

EDC (540mg, 3mmol) was added to a stirred solution of (O-2,6dichlorobenzyl)-L-tyrosine methyl ester (1g, 2.6mmol), 2 methylmercaptonicotinic acid (433mg, 2.6mmol), HOBT (364mg, 2.6mmol) and NMM (846ul, 7.8mmol) in anhydrous DMF (4ml) at 0°. The DMF solution was stirred overnight at room temperature, then the DMF was evaporated in vacuo. The residue was taken up in DCM (70ml), washed with water (3 x 15ml), saturated aqueous NaHCO3 (2 x 15ml) and water (2 x 15ml), dried (NaoSO_A) and evaporated in vacuo. The residue was purified by chromatography (SiO2; EtOAc) to give the title compound as a white solid (1.2gm, 92%). SH (CDCl₃) 8.42 (1H, dd, <u>J</u> 1.7, 4.8, 1pyrH)(, 7.67 (1H, dd, J 1.7, 7.6, 1pvrH), 7.31 (2H, d, J 8.5, ArH), 7.18 (1H, dd, J 2.3, 8.5, ArH), 7.09 (3H. d. plus broad peak, J 8.6, 2ArH, 1NH), 6.97 (1H, m, 1pyrH), 6.88 (2H, d, J 8.6, 2ArH), 5.18 (2H, s, OCH2Ar), 4.96 (1H, m, CHatyr), 3.72 (3H, s, OCH₃), 3.25-3.04 (2H, m, CH₂Ar) and 2.47 (3H, s, SCH₃). m/z (ESI, GOU) 505 (MH+).

INTERMEDIATE 11

Ethyl 3-(4-(((4-methoxybenzyl)oxy)carbonyl)phenyl)-2-

[(diphenylmethylene)amino)propanoate

N-(Diphenylmethylene) glycine ethyl ester (6.6g, 24.6mmol) and potassium carbonate (6.8g, 49mmol) were added to a solution of 4-methoxybenzyl-4(bromomethyl)benzoate (8.2g, 24.6mmol) in acetonitrile (200ml). The mixture was heated at reflux ovenight, then filtered and the solvent removed *in vacuo* to give the <u>title compound</u> as a yellow oil (13.55g). δ H (CDCl₃, 300 MHz) 7.8 (2H, d, $\frac{1}{2}$ 9.0Hz), 7.5 (10H, m), 7.3 (2H, d, $\frac{1}{2}$ 9.0Hz), 6.9 (2H, d), 6.6 (2H, m), 5.23 (2H, s), 4.1 (3H, m), 3.7 (3H, s), 3.2 (2H, m) and 1.2 (3H, m); $\frac{mv}{mv}$ (ESI) 522 (MH+).

INTERMEDIATE 12

Ethyl 2-amino-3-(4[[(4-methoxybenzyl)oxy]carbonyl)phenyl)

10 propanoate

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Hydrochloric acid (2M, 15.83ml, 1.5eq) was added to a solution of Intermediate 11 (11.0g, 21.12mmol) in THF (30ml). After 20min the reaction mixture was basified to pH7 with NaHCO₃ and the solvent removed *in vacuo*. The residue was taken up in EtOAc (300ml) and washed with water (200ml) and brine (200ml), dried (MgSO₄) and evaporated *in vacuo*. Chromatography (SiO₂; EtOAc) gave the <u>title compound</u> as a yellow oil (4.91g, 65%). δ H (CDCl₃, 300MHz) 7.88 (2H, d. \pm 8.0Hz), 7.4 (4H, dd), 7.0 (2H, d, \pm 8.0Hz), 5.25 (2H, s), 4.05 (2H, q), 3.7 (3H, s), 3.57 (1H, t), 2.87 (2H, m) and 1.1 (3H, t), *m/z* (ESI) 358 (MH+).

INTERMEDIATE 13

Ethyl-2-[[(2-chloro-3-pyridinyl)carbonyl]amino]-3-(4-[[(4-

methoxybenzyl)oxylcarbonyl)phenyl)propanoate

EDC;HCI (591 mg, 3.08mmol) and HOBT (416 mg, 3.08 mmol) were added to a solution of intermediate 12 (1.0g, 2.8 mmol), 2-chloronicotinic acid (450 mg, 2.86 mmol) and NMM (370 µl, 3.36 mmol) in DMF (30 ml). The reaction mixture was stirred overnight at room temperature. The solvent was removed in vacuo and the residue partitioned between EtOAc (300 ml) and NaHCO3 solution (300 ml). The organic phase was washed with citric acid (10%, 2 x 200 ml), NHCO3 solution (200 ml) and brine (300 ml), dried (MgSO4) and concentrated in vacuo to give the title compound as a yellow oil (1.36g, 98%). &H (CDCl3, 300 MHz) 9.12 (1H, d, ½ 8.0 Hz), 8.47 (1H, m), 7.91 (2H, d, ½ 8.0 Hz), 7.6 (1H, dd), 7.45 (5H, m), 6.95 (2H, d, ½ 8.0 Hz), 4.67 (1H, m), 4.15 (2H, m), 3.75 (3H, s), 3.31 (1H, m), 3.23 (1H, m) and 1.17 (3H, s); m/z (ESI) 497 (MH+).

INTERMEDIATE 14

Ethyl 2-([(2-chloro-3-pyridinyl)carbonyl]amino)-3-[4-(carboxyl)phenyl] propanoate

TFA (20ml) was added to a solution of Intermediate 13 (1.36g, 2.75mmol) in toluene (20ml). The reaction mixture was stirred for 30min at room temperature. The white solid obtained was recrystallised (EtOAc/hexane) to give the <u>title compound</u> (1.04g, 100%). &H (CDCl₃, 300MHz) 12.84 (1H, br s), 9.1 (1H, d, <u>J</u> 8.0Hz), 8.47 (1H, m), 7.87 (2H, d, <u>J</u> 8.0HZ), 4.1 (2H, m), 3.2 (1H, m), 3.07 (1H, m) and 1.17 (3H, m); m/z (ESI) 377 (MH*).

INTERMEDIATE 15

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Ethyl 2-{[(2-chloro-3-pyridinyl)carbonyl]amino}-3-(4-{(2.6-d)chloroanilino)carbonyl}phenyl)propanoate

Carbon tetrachloride (5ml) was added to a suspension of Intermediate 14 (1.04g, 2.78mol) and triphenylphosphine (0.87g, 3.31mmol) in acetontrile. The reaction mixture was stirred for 2h at room temperature. 2,6-Dichloroaniline (0.89g, 5.52mmol) and NMM (455µl, 4.14mmol) were added and the mixture stirred for a further 48h at room temperature. The solvent was removed *in vacuo* and the residue partitioned between water and EtOAc. The aqueous layer was extracted with EtOAc (x 2) and the combined organic extracts washed with water (x 2) and saturated aqueous NaHCO₃ (x2), dried (Na₂ SO₄) and concentrated *in vacuo*. Chromatography (SiO₂; EtOAc/hexane 50:50) gave the title compound (924mg) 8H (CDCl₃, 300MHz) 8.46 (1H, s), 8.05 (1H, d, ½ 7.4Hz), 7.89 (2H, d, ½ 7.9Hz), 7.69-7.10 (8H, m), 5.11 (1H, dt, ½ 6.7, 5.8Hz), 4.25 (2H, q, ½ 7.1Hz), 3.42 (1H, dd, ½ 13.9, 5.8Hz), 3.31 (1H, dd, ½ 13.9, 5.8Hz) and 1.30 (3H, t, ½ 7.1Hz).

INTERMEDIATE 16

Methyl 4-[2/2.6-dlchlorophenyl)-2-hydroxylethyl]benzoate

A solution of methyl 4-(bromomethyl)benzoate (2.0g, 8.7mmol) in THF (4.4ml) was added slowly to cut zinc foil (683mg, 10.44mmol) which had been activated with 1,2-dibromoethane (80mg). After 3h of stirring at room temperature 2ml of the solution was transferred to a solution of copper cyanide (396mg, 4.4mmol) and lithium chloride (356mg, 8.4mmol) in THF (4ml) cooled to -78°. This solution was warmed to -20° and then cooled

back to -78°. Boron trifluoride etherate (983µl, 8mmol) was then added followed by 2,6-dichlorobenzaldehyde (0.56g, 3.2mmol) in THF (1ml). The reaction was stirred for 2h and then allowed to warm slowly to room temperature. Water (20ml) was then added and the reaction mixture extracted into EtOAc (3 x 25ml) and the combined organics dired (Na₂SO₄) and evaporated. Purification by column chromatography (SiO₂; hexane:EtOAc. 4:1) gave the title compound as a colourless oil (863mg, 83%). 8H (CDCl₃) 7.95 (2H, m, ArH), 7.93-7.26 (4H, m, ArH), 7.17-7.12 (1H, m, ArH), 5.73-5.65 (1H, m, CH), 3.91 (3H, s, CO₂Me), 3.43 (1H, dd, ½ 13.5, 8.4Hz, CH_AH_B) and 3.28 (1H, dd, ½ 13.5, 6.3Hz, CH_AH_B), m/z (ESI, 60V) 325 (MH+).

INTERMEDIATE 17

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Methyl 4-[2-[(tert-butyl(dimethyl)silyl]oxy)-2-(2.6-dichlorophenyl) ethyl]benzoate

To a solution of Intermediate 16 (20g, 6.15mmol) in DCM (10ml) cooled to 0° was added 2,4,6-collidine (2.03ml, 15.39mmol). After 15min terbutyldimethylsilyltrifluoromethanesulphonate (2.12ml, 9.23mmol) was added. The reaction mixture was stirred overnight at room temperature then diluted with DCM (100ml) and washed with 1M hydrochloric acid (50ml), water (50ml) and brine (50ml), dried (Na₂SO₄) and evaporated under reduced pressure. Purification by column chromatography (SiO₂; hexane:EtOAc, 5:1) gave the title compound as a pale pink oil (2.67g, 100%). 8H (CDCl₃) 7.94 (2H, d, ½ 6.5Hz, ArH), 7.33, 7.21 (4H, m, ArH), 7.13-7.07 (1H, m, ArH), 5.58 (1H, dd, ½ 9.4, 4.6Hz, CH), 3.90 (1H, s, CO₂Me), 3.46 (1H, dd, ½ 13.1, 9.4Hz, CH₂H₃), 3.04 (1H, dd, ½ 13.1, 4.6Hz, CH₃H₃), 0.74 (9H, s, Si¹Bu), -0.31 (3H, s, SiMe) and -0.32 (3H, s, SiMe); m²z (ESI, 60V) 361 (MH+).

30 INTERMEDIATE 18

4-[2-([Tert-butyl(dimethyl)silyl]oxy]-2-(2.6-

dichlorophenyl)ethyl]benzylalcohol

Lithium aluminium hydride (1M solution in THF, 6.46ml, 6.46mmol) was added to an ice cold solution of Intermediate 17 (2.67g, 6.15mmol) in THF (20ml). The reaction mixture was stirred for 1h then quenched with the addition of water and extracted into DCM (3 x 50ml), dried (Na₂SO₄) and

evaporated under reduced pressure. Purification by column chromatography (SiO₂; hexane:EtOAc, 4:1) gave the <u>title compound</u> as a colourless oil (2.g, 87%) δH 7.32-7.06 (7H, m, ArH), 5.56 (1H, dd, <u>J</u> 9.3, 4.7Hz, CH₂C<u>H</u>), 4.65 (2H, d, <u>J</u> 5.9Hz, C<u>H</u>₂C), 3.39 (1H, dd, <u>J</u> 13.2, 9.3Hz, CH_AH_B), 3.01 (1H, dd, <u>J</u> 13.2, 4.7Hz, CH_AH_B), 0.75 (9H, s, Si\(\frac{B}{2}\)u), -0.29 (3H, s, Si\(\frac{M}{2}\)u) and -0.31 (3H, s, Si\(\frac{M}{2}\)u) in (2 (ESI, 60V) 433 (MH+).

INTERMEDIATE 19

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4-[2-([Tert-butyl(dimethyl)silyl]oxy]-2-(2,6-dichlorophenyl)ethyl] benzylbromide

A solution of triphenylphosphine (843mg, 3.21mmol) in DCM (2ml) was added to a solution of carbon tetrabromide (1.42g, 3.73mmol) and Intermediate 18 (1.10g, 2.67mmol) in DCM (3ml) and stirred at room temperature for 24h. Ether (100ml) was added and the solid precipitate formed removed by filtration. The filtrate was evaporated under reduced pressure and the residue purified by column chromatography (SiO₂: 8:1, hexane:EtOAc) to give the <u>fitle compound</u> (1.20g, 95%). δH (CDCl₃) 7.32-7.07 (7H, m, ArH0, 5.54 (1H, dd, ½ 9.5, 4.4Hz, CH₂CH₃), 4.49 (2H, s. CH₂Br), 3.39 (1H, dd, ½ 13.2, 9.5Hz, CH₂H_B), 2.96 (1H, dd, ½ 13.2, 4.4Hz, CH₄H_B), 0.73 (9H, s. SiN₂D₃), -0.31 (3H, s. SiM₂D) and -0.32 (3H, s. SiM₂D), y(z (ESI, 60V) 474 (MH+).

INTERMEDIATE 20

4-[2-([Tert-butyl(dimethyl)silyl]oxy]-2-(2.6-dichlorophenyl)ethyl] phenylalanine ethyl ester

To a solution of ethyl *N*-(diphenylmethylene)glycinate (2.63g, 9.81mmol) in THF (50ml) cooled to -78° was added lithium diisopropylamine (2M in heptane/THF/ethylbenzene, 5.64ml, 11.28mmol). The solution was stirred for 45min. Intermediate 19 (4.20g, 8.92mmol) in THF (20ml) was then added dropwise. The reaction mixture was stirred for 2h at -78° and then warmed to room temperature. EtOAc (100ml) was added and the mixture washed with water (75ml) and brine (75ml), dried (Na₂SO₄) and evaporated under reduced pressure. The residue was then taken up in acidic ethanol and stirred for 10min. The volatiles were then removed and the residue partitioned between EtOAc (150ml) and saturated aqueous Na₂CO₃ (100ml). The aqueous layer was extracted several times with

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EtOAc and the combined organics dried (Na₂SO₄) and evaporated under reduced pressure. The remaining residue was purified by column chromatography (SiO₂; EtOAc) to give the <u>title compound</u> as a colourless oil (3.95g, 95%). 8H (CDCl₃) 7.32-7.05 (7H, m, ArH), 5.52 (1H, dd, $\frac{1}{2}$ 9.4, 4.6Hz, CHOSl), 4.18 (2H, q, $\frac{1}{2}$ 7.1Hz, CO₂CH₂CH₃), 3.68 (1H, dd, $\frac{1}{2}$ 7.9, 5.1Hz, CHNH₂), 3.35 (1H, dd, $\frac{1}{2}$ 13.2, 9.4Hz, CHOSiCH₂H₃), 3.07 (1H, dd, $\frac{1}{2}$ 13.5, 5.1Hz, CHNH₂CH₂H₃), 2.95 (1H, dd, $\frac{1}{2}$ 13.2, 4.6Hz, CHOSiCH₄H₃), 2.95 (1H, dd, $\frac{1}{2}$ 13.5, 7.9Hz, CHNH₂CH₄H₃), 1.53 (2H, br s, NH₂), 1.27 (3H, t, $\frac{1}{2}$ 7.1Hz, CO₂CH₂CH₃), 0.74 (9H, s, SiBu) and -0.32 (6H, s, SiMe₂); m/z (ESI, 60V) 496 (MH+).

INTERMEDIATE 21

[4-[2-[[Tert-butyl(dimethyl)silyl]oxy]-2-(2,6-dichlorophenyl)ethyl]]-(N-2-chloronicotinoyl)phenylalanine ethyl ester

To a solution of Intermediate 20 (1.50g, 3.2ommol) and 2-chloronicotinic acid (504mg, 3.20mmol) in DCM (75ml) at room temperature was added NMM (386µI,3.53mmol), EDC (675mg, 3.53mmol) and HOBT (477mg, 3.53mmol). The reaction mixture was stirred overnight at room temperature and then diluted with DCM (50ml) and washed with saturated aqueous Na₂CO₃ (50ml), water (50ml) and brine (50ml), dried (Na₂SO₄) and evaporated under reduced pressure. The residue was purified by column chromatography (SiO₂; EtOAc) to give the title compound as a white solid (1.72g, 85%). &H (CDCl₃) &.41-8.38 (1H, m, NH), 8.01-7.96 (1H, m, ArH), 7.29-6.97 (9H, m, ArH), 5.47 (1H, dd, 1.9.2, 4.7Hz, CHOSi), 5.24-4.96 (1H, m, CHNH), 4.19 (2H, qd, 1.7.1, 1.1Hz, CO₂CH₂CH₃), 3.34-3.14 (4H, m, CH₂ x 2), 1.27 (3H, td, 1.7.1, 1.3Hz, CO₂CH₂CH₃), 0.69 (s) and 0.66 (s); together (9H, SitBu) and -0.38 (s) and -0.40 (s); together (6H, SitBo); 659 (M*+ Na*).

INTERMEDIATE 22

(N-2-Chloronicotinoyi)-4-[2-(2.6-dichlorophenyi)-2-hydroxyethyl) phenylalanine ethyl ester

Tetrabutylammonium fluoride (1M in THF, 4.7ml, 4.70mmol) was added to a solution of Intermediate 21 (1.50g, 2.35mmol) in THF (75ml) at room temperature. The reaction mixture was stirred for 2h and then the THF

removed and the residue partitioned between EtOAc and water. The layers were separated and the aqueous layer extracted with EtOAc. The combined organics were dried (Na₂SO₄) and evaporated under reduced pressure. Purification of the residue by column chromatography (SiO₂; MeOH:DCM, 5:95) gave the <u>title compound</u> as a pale brown oil (1.03g, 84%) δH (CDCl₃) 8.31-8.29 (1H, m, NH), 7.86-7.82 (1H, m, ArH), 7.25-6.96 (9H, m, ArH), 5.54-5.49 (1H, m, CH), 4.94-4.88 (1H, m, CH), 4.11 (2H, qd, <u>1</u> 7.1, 2.0Hz, CO₂CH₂CH₃); m/z (ESI, 60V) 521 (MH+).

INTERMEDIATE 23

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(N-2-Chloronicotinoyl)-{4-[2-(2.6-dichlorophenyl)-2-oxoethyl]} phenylalanine ethyl ester

To a solution of Intermediate 22 (300mg, 0.58mmol) in acetone (20ml) was added Jones' Reagent dropwise until an orange colour persisted. i-Propyl alcohol was added to use up excess reagent and then the solution was basified by the addition of saturated aqueous Na₂CO₃ solution. The solution was then decanted from the solids and the acetone removed in vacuo. The remaining aqueous solution was then extracted with ether (x 2) and the combined organics dried (Na₂SO₄) and evaporated under reduced pressure. The residue was purified by column chromatography (SiO₂; hexane:EtOAc, 3:2) to give the title compound as a colourless oil (200mg, 67%). &H (CDCl₃) 8.40 (1H, dd, 1 4.8, 2.0Hz), 7.94 (1H, dd, 1 7.7, 2.0Hz), 7.38-7.11 (8H, m, ArH), 7.02 (1H, d, 1 7.5Hz, ArH), 4.19 (2H, q, 1 7.2Hz, CO₂CH₂CH₃), 4.09 (2H, s, CH₂CO), 3.28 (1H, dd, 1 14.0, 5.9Hz, CH₃H₃), 3.18 (1H, dd, 1 14.0, 6.1Hz, CH₃H₃) and 1.26 (5H, t, 1 7.2Hz, CO₂CH₂CH₃); m/z (ESI, 60V) 519 (MH+*).

INTERMEDIATE 24

Methyl 4-f(E)-2-(2,6-dichlorophenyl)ethenylibenzoate

A solution of Intermediate 16 (2.0g, 6.15mmol) in toluene (25ml) containing p-toluenesulphonic acid (100mg) was heated to reflux in a Dean-Stark apparatus for 4h. Toluene was then removed under reduced pressure and the residue purified by column chromatography (SiO₂; hexane:EtOAc, 5:1) to give the title compound as an off white solid (1.64g;87%); 8H (CDCl₃) 8.05 (2H, d, <u>J</u> 8.3Hz), 7.60 (2H, d, <u>J</u> 8.3Hz), 7.36 (2H, d, <u>J</u> 8.0Hz),

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7.21 (2H, d, \downarrow 1.7Hz), 7.15-7.10 (1H, m), and 3.93 (3H, s, CO₂CH₃); <u>m/z</u> (ESI, 60V) 329 (MH+).

INTERMEDIATE 25

4-[(E)-2-(2,6-Dichlorophenyl)ethenyl]benzyl alcohol

To an ice cold solution of Intermediate 24 (1.56g, 5.08mmol) in THF (20ml) was added lithium aluminium hydride (1M inTHF, 5.34ml, 5.34mmol). The reaction mixture was stirred for 30min and then quenched by the addition of water (10ml). The resulting biphasic solution was filtered through Celite® and then extracted with DCM (2 x 50ml). The combined organics were dried (Na₂SO₄) and evaporated under reduced pressure to give the title compound as a colourless oil which solidified on standing (1.5g, 99%). 8H (CDCl₃) 7.55 (2H, d, J 8.2Hz), 7.42-7.33 (4H, m), 7.20-7.08 (3H, m) and 4.75 (2H, d, J 5.3Hz, CH₂); m/z (ESI, 60V) 301 (M*+ Na*+).

INTERMEDIATE 26

4-f(E)-2-(2.6-Dichlorophenyl)ethenyl]benzyl bromide

A solution of triphenylphosphine (1.58h, 6.04mmol) in DCM (10ml) was added to a solution of intermediate 25 (1.40g, 5.03mmol) and carbon tetrabromide (2.33g, 7.04mmol) in DCM (10ml). The resulting solution was stirred for 1h and then diluted with ether (150ml) and the resulting solid removed by filtration. The filtrate was then evaporated under reduced pressure and the resulting residue purified by column chromatography (SiO₂; hexane: EtOAc, 6:1) to give the title compound as a colouless oil (1g, 49%). 8H (CDCb) 7.55 (2H, d, J 8.3Hz), 7.44-7.32 (3H, m), 7.23-7.09 (4H, m) and 4.53 (2H, s, CH₂); m/z (ESI, 60V) 342 (MH+)

INTERMEDIATE 27

4-[(E)-2-(2.6-Dichlorophenyl)ethenyl]phenylalanine ethyl ester

30 To a solution of ethyl A-(diphenylmethylene)glycinate (860mg,3.21mmol) in THF (20ml) cooled to -78° was added lithium diisopropylamine (2M in heptane/THF/ethylbenzene, 1.68ml, 3.36mmol). The solution was stirred for 45min. Intermediate 26 (10g, 2.92mmol) in THF (2ml) was added and the resulting reaction mixture stirred for 2h at -78° and then warmed to room temperature. EtOAc (100ml) was added and the mixture washed with water (75ml) and brine (75ml), dried (Na₂SO₄) and evaporated under

reduced pressure. The residue was then taken up in acidic ethanol and stirred for 1min. The volatiles were then removed and the residue partitioned between EtOAc (150ml) and saturated aqueous Na₂CO₃ (100ml). The aqueous layer was extracted several times with EtOAc and the combined organics dried (Na₂SO₄) and evaporated under reduced pressure. The remaining residue was purified by column chromatography (SiO₂; EtOAc) to give the title compound as a colouress oil (860mg, 81%). 5H (CDCl₃) 7.47 (2H, d, J. 8.1Hz), 7.32 (2H, d, J. 8.1Hz), 7.21 (2H, d, J. 8.2Hz), 7.19-6.98 (3H, m), 4.18 (2H, q, J. 7.1Hz, CO₂CH₂CH₃), 3.71 (1H, dd, J. 7.8, 5.3Hz, CH), 3.10 (1H, dd, J. 13.5, 5.3Hz, CH₃H₃), 2.88 (1H, dd, J. 13.5, 7.8Hz, CH₃H₃B) and 1.26 (3H, t, J. 7.1Hz, CO₂CH₂CH₃); m/z (ESI, 60V) 364 (MH+)

INTERMEDIATE 28

(N-2-Chloronicotinoyl) 4-[(5)-2-(2.6-dichlorophenyl)ethenyl] phenylalanine ethyl ester

To a solution of intermediate 27 (860mg, 2.36mmol) and 2-chloronicotinic acid (372mg, 2.36mmol) in DCM (25ml) was added NMM (285μl, 2.60mmol), EDC (498mg, 2.60mmol) and HOBT (352mg, 2.60mmol). The resulting solution was stirred for 3h and then diluted with DCM (50ml) and brine (50ml), dried (Na₂SO₄) and evaporated under reduced pressure. The residue was purified by column chromatography (SiO₂; EtOAc) to give the title compound as a pale yellow oil (1.1g, 93%). δH (CDCl₃), 8.43 (1H, dd, ½ 4.8, 2.0Hz), 8.02 (1H, dd, ½ 7.6, 2.0Hz), 7.46 (2H, d, ½ 8.2Hz), 7.34-6.92 (9H, m), 5.10-5.03 (1H, m, CH),4.24 (2H, q, ½ 7.1Hz, CO₂CH₂CH₃), 3.33 (1H, dd, ½ 13.9, 5.8Hz, CHAH_B), 3.24 (1H, dd, ½ 13.9, 5.8Hz, CHAH_B) and 1.30 (3H, t, ½ 7.1Hz, CO₂CH₂CH₃).

INTERMEDIATE 29

N-(2-Chloronicotinovi)-L-tyrosine methyl ester

EDC.HCI (2.11g, 11mmol) was added to a mixture of *L*-tyrosine methyl ester hydrochloride (2.32g, 10mmol), 2-chloronicotinic acid (1.58g, 10mmol), HOBT (1.49g, 11mmol) and NMM (2.31ml, 21mmol) in DMF (50ml). The mixture was stirred overnight at room temperature. The solvent was removed *in vacuo*. the residue dissolved in EtOAc (300ml) and washed with dilute HCI (100ml), saturated aqueous NaHCO₃

(100ml), water (3 x 100ml) and brine (50ml), dried (NaoSO₄) and solvent removed in vacuo to give the title compound as a vellow gum (3.27g. 98%), δH (DMSO-de, 300MHz) 9.21 (1H, s, OH), 9.03 (1H, d, J 7.9Hz, CONH), 8.45 (1H, dd, J 4.8, 1.9Hz, pyrH), 7.67 (1H, dd, J 7.4, 1.9Hz, pyrH), 7.47 (1H, dd, J 7.5, 4.8Hz, pyrH), 7.05 (2H, d, J 8.5Hz, ArH), 6.67 (2H, d, <u>J</u> 8.5Hz, ArH), 4.58 (1H, ddd, <u>J</u> 9.6, 7.9, 5.4Hz, CHα), 3.65 (3H, s, CO₂Me), 3.01 (1H, dd, J 13.9, 5.4Hz, CH_AH_BAr) and 2.85 (1H, dd, J 13.9, 9.6Hz, CHAHBAr), m/z (ESI, 60V) 335 (MH+).

10 INTERMEDIATE 30

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N-(2-Chloronicotinovi)- O-(2.6-dichlorobenzovi)- L-tyrosine methyl

A solution of Intermediate 29 (919mg, 2.75mmol) in DMF (10ml) was added to a suspension of sodium hydride (60% in mineral oil, 3.03mmol, 121mg) in DMF (20ml) at 0°. After 15min, 2.6-dichlorobenzoyl chioride (414µl, 2.89mmol) was added and the mixture stirred for 2h at room temperature. Water (~5ml) was added and the solvent removed in vacuo. The residue was dissolved i nEtOAc (150ml), washed with water (3 x 50ml) and brine (50ml), dried (Nao SO₄) and evaporated in vacuo. Chromatography (SiO2: DCM/MeOH, 98:2) gave the title comound as a 20 white foam (1.10g, 79%). δH (DMSO-d₆, 300MHz) 9.12 (1H, d, J 7.6Hz, CONH), 8.45 (1H, dd, J 4.8, 2.0Hz, pyrH), 7.69-7.58 (4H, m, pyrH + Cl₂ArH₃), 7.46 (1H, dd, <u>J</u> 7.5, 4.9Hz, pyrH), 7.43 (2H, d, <u>J</u> 8.4Hz, ArH₂), 7.23 (2H, d, <u>J</u> 8.5Hz, ArH₂), 4.73 (1H, m, CHα), 3.68 (3H, s, CO₂Me), 3.22 (1H, dd, <u>J</u> 13.9, 5.2Hz, CHAHBAr) and 3.02 (1H, dd, <u>J</u> 13.9, 10.1Hz, 25 CHAHBAr): m/z (ESI, 60V) 507 (MH+).

INTERMEDIATE 31

N-(2-Chloronicotinovi)-N-methyl-N'-(3.5-dichloro-4-picolyl)-L-4aminophenvialanine methyl ester

2-Chloronicotinoyl chloride (132mg, 0.75mmol) was added to N-methyl-N'-(3,5-dichloroisonicotinyl)-L-4-aminophenylalanine methyl ester [prepared from N-Boc-N'-phthaloyl-4-amino-L-phenylalanine methyl ester and methyl iodide, followed by treatment with hydazine monohydrate and reaction with 3.5-dichloroisonicotinyl choride with subsequent removal of the Boc group] and NMM (165µl, 1.5mmol) in DCM (10ml). The mixture was stirred for 1h at room temeprature then diluted with DCM (100ml) and washed with dilute HCI (30ml), dried (Na₂SO₄) and evaporated *in vacuo*. Chromatography (SiO₂; EtOAc/hexane, 10:90) gave the <u>titlé compound</u> as a colourless gum (380mg, 97%). δ H (DMSO-d₆, 300MHz, 405K) 10.35 (1H, br s, CONH), 8.67 (2H, s, Cl₂pyrH), 8.42 (1H, t, \pm 3.4Hz, ClpyrH), 7.55 (2H, br d, \pm 7.2Hz, ArH), 7.45-7.15 (4H, v br m, ArH + ClpyrH), 5.3 (1H, v br s, CHa) , 3.74 (3H, s, CO₂Me), 3.4-3.3 (1H, br m, CHaHBAr), 3.16 (1H, dd, \pm 14.4, 9.6Hz, CHaHBAr) and 2.73 (3H, br s, NMe): m/z (ESI, 60V) 521 (MH+).

INTERMEDIATE 32

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[(S-2,5-dlmethoxyphenyi)sulphonyi]nicotinoyl-O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester

A solution of Intermediate 2 (0.79g, 1.26mmol) in DCM (50ml) was treated with 3-chloroperoxybenzoic acid (2.17g, 12.6mmol) and stored at 4° for 48h. The reaction was partitioned between DCM (20ml) and NaHCO₃ solution (20ml). The aqueous layer was extracted with DCM (25ml) and the combined organic layers washed with 10% aqueous Na₂SO₃ (50ml), dried (MgSO₄) and the solvent removed *in vacuo* to give a yellow foam that was purified by chromatography (SiO₂; EtOAc/hexane 3:1) to give the title compound as a white foam, (0.50g, 60%). 8H (CDCl₃) 8.58 (1H, dd, 1.4.7, 1.7Hz, pyr-H), 7.95 (1H, dd, 1.7.8, 1.7Hz, pyr-H), 7.70 (1H, d, 1.32Hz, Ar-H), 7.50 (1H, dd, 1.7.8, 4.7Hz, pyr-H), 7.36-7.11 (6H, m. Ar-H), 7.01 (1H, d, 1.7.6Hz, NH), 6.92-6.84 (3H, m, Ar-H), 5.21 (2H, s, CH₂O), 5.10 (1H, m, CHc₃), 3.85 (3H, s, OMe), 3.74 (3H, s, OMe), 3.49 (3H, s, CO₃Me) and 3.27 (2H, m, CHCH₂An), myz (ESI, 60V) 659 (MH+).

EXAMPLE 1

2-Chloronicotinovi-O-(2,6-dichlorobenzyl)-L-tyrosine

A solution of Intermediate 1 (0.20g, 0.41mmol) in THF (5ml) and water (5ml) was treated with lithium hydroxide monohydrate (25ml, 0.61mmol. 1.5 equiv.) and stirred at room temperature for 1.5h. The reaction was acidified to pH1 with 10% hydrochloric acid to give a white precipitate which was isloated by filtration, washed with water (5ml) and died under vacuum to give the title compound as a white

powder (0.14g, 72%). δ H (DMSO-d⁶) 8.93 (1H, d, \downarrow 8.1Hz), 8.45 (1H, dd, \downarrow 2.0, 4.8Hz), 7.66 (1H, dd, \downarrow 2.0, 7.5Hz), 7.56 (2H, m), 7.47 (2H, m), 7.24 (2H, d, \downarrow 8.6Hz), 6.99 (2H, d, \downarrow 8.6Hz), 5.20 (2H, s), 4.60 (1H, d), 3.13 (1H, ABX, \downarrow 4.7, 13.9Hz) and 2.90 (1H, ABX, \downarrow 10.0, 13.9Hz).m/z (ES+, 60V) 479, 481 (MH+).

EXAMPLE 2

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2-Thio(S-2,5-dimethoxyphenyl)nicotinoyl-O-(2,6-dicholorobenzyl)-L-tyrosine hydrochloride

A solution of Intermediate 2 (0.35g, 0.56mmol) in THF (15ml) and water (7.5ml) was treated with lithium hydroxide monohydrate (28mg, 0.67mmol) and stirred at room temperature for 16h. The reaction was acidified to pH1 with 10% hydrochloric acid, extracted with DCM (2 x 30ml), and the combined organic layers were dried over MgSO₄, the solvent removed in vacuo to give a gummy residue, which was purified by chromatography (SiO₂; 7.5% MeOH/DCM) to give a gum which was dissolved in acetonitrile (20ml) and water (20ml) and lyophylised to give the title compound as a white powder (0.26g, 71%). 8H (DMSO-d9) 8.71 (1H, m), 8.29 (1H, dd, J. 1.7, 4.8Hz), 7.75 (1H, d, J. 6.0Hz), 7.56-6.94 (11H, m), 5.16 (2H, s), 4.56 (1H, m), 3.68 (3H, s), 3.60 (3H, s) and 3.21-1.96 (2H, m). m/z (ES+60V) 613, 615 (MH+).

EXAMPLE 3

a) 2-Thio(S-4-picolinyl)nicotinovi-O-(2.6-dichlorobenzyl)-L-tyrosine

A solution of Intermediate 4 (0.30g, 0.52mmol) in THF (7.5ml) and water (5ml) was treated with lithium hydroxide monohydrate (33mg, 0.7mmol) and stirred at room temperature for 16h. The pH was adjusted to 6.5-7 with 10% hydrochloric acid to give a yellow precipitate which was isolated by filtration, washed with water and dried *in vacuo* to give the title compound as a yellow powder (0.28g, 95%). 8H (DMSO-d6) 8.83 (1H, d, $\frac{1}{2}$ 8.9Hz), 8.5 (3H, m), 7.75 (1H, d, $\frac{1}{2}$ 7.6Hz), 7.56-7.20 (8H, m), 6.95 (2H, d, $\frac{1}{2}$ 8.5Hz), 5.17 (2H, s), 4.54 (1H, m), 4.37 (2H, s), 3.12 (1H, m) and 2.97 (1H, m), m/z (ES+, 60V) 568, 570 (MH+).

The following compouns were prepared in a similar manner by hydrolysis of the corresponding methyl ester. Each ester starting material was obtained either by alkylation of intermediate 3 or alterntive

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mercaptopyridine using the reagents shown using a similar procedure to that described for intermediate 4:

b) 2-Thio-S-benzyl-nicotinoyl-(0-2.6-dichlorobenzyl)-L-tyrosine ester from intermediate 3 and benzyl chloride. δH (DMSO-d₆) 8.78 (1H, d, \downarrow 8.1Hz, pyr-H), 8.54 (1H, dd, \downarrow 4.8, 1.Hz, pyr-H), 7.69 (1H, dd, \downarrow 7.7,1.7Hz, pyr-H), 7.68-7.17 (11H, m, Ar-H), 6.94 (2H, d, \downarrow 8.6, Ar-H), 5.16 (2H, s, CH₂O), 4.53 (1H, m, CH α), 4.34 (2H, s, CH₂Ar), 3.10 (1H, dd, \downarrow 13.9, 4.6Hz, CHCH_AH_BAr); m/z (ESI, 60V) 567 (MH+).

c) 2-Thio-(S-4-Methylphenyl)-nicotinoyi-(0-2.6-dichlorobenzyl-L-tyrosine

d) 2-Thio-S-(3-picolyi)-nicotinoyi-(0-2,6-dichlorobenzyi)-L-tyrosine ester from intermediate 3 and 3-picolyi chloride. 8H (DMSO-de) 9.04-8.54 (5H, m, Ar-H), 7.95-7.84 (2H, m, Ar-H), 7.56-7.43 (3H, m, Ar-H), 6.96 (2H, d, <u>J. 8.3Hz</u>, Ar-H), 5.17 (2H, s, CH₂O), 4.55 (1H, m, CHa₂), 4.48 (2H, s, CH₂pyr), 3.05 (2H, m, CHC<u>H</u>₂Ar); m/z (ESI, 60V) 559 (MH*).

e) <u>N-[2-thio(S-3-picolinyl)nlcotinoyl]-0-,2.6-dichlorobenzyl-L-</u>tyrosine

ester from Intermediate 3 and 3-picoyl chloride using DBU as base. δH (DMSO-d₆) 8.8 (1H, br d), 8.57 (1H, m), 8.50 (1H, m), 7.72 (2H, m), 7.55 (2H, d), 7.5-7.4 (2H, m), 7.3-7.15 (4H, m), 6.95 (2H, d), 5.15 (2H, s), 4.55-4.45 (3H, m), 3.2-3.1 (1H, m), 3.0-2.9 (1H, m); m/χ (ESI, 60V) 588 (MH+γ).

f) N-2-Thio(S-4-butanoate)nicotinoyi]-(O-2,8-dichlorobenzyi)-L-tyrosine

ester from Intermediate 3 and methyl-4-chlorobutyrate using K_2CO_3 as base. δH (DMSO-d₆) 8.76 (1H, d, \downarrow 8.1Hz), 8.49 (1H, dd, \downarrow 4.8, 1.7Hz), 7.61 (1H, dd, \downarrow 7.7, 1.7Hz), 7.55 (2H, d, \downarrow 8.9Hz), 7.45 (1H, m), 7.23 (2H, d, \downarrow 8.5Hz), 7.2 (1H, m), 6.96 (2H, d, \downarrow 8.5Hz), 5.18 (2H, s), 4.53 (1H, m), 3.17-2.87 (4H, m), 2.40-2.30 (2H, t, \downarrow 7.3Hz), 1.83 (2H, m); \underline{m}/z (ESI, 60V) 563 (MH+).

g) (N'-3.5-Dichloroisonicotinoyi)-N-(([3-pyridinylmethyl]thio) isonicotinoyi)-L-4-aminophenylalanine

ester from 2-mercaptoisonicotinoyi-(N'-3,5-dichloroisonicotinoyi)-L-4-aminophenyialanine and 3-picolyl chloride. 8H (DMSO-de) 10.86 (1H, s, CO₂H), 8.92-8.75 (3H, m, ArH), 8.68-8.52 (2H, m), 8.39 (1H, br s), 7.75 (2H, t, <u>J</u> 7.5Hz), 7.56 (2H, d, <u>J</u> 6.3Hz), 7.39-7.18 (4H, m), 4.60-4.47 (1H, m, CH) and 3.25-2.92 (2H, m, CH₂); m/z (ESI, 60V) 582 (MH+).

EXAMPLE 4

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a) 2-Thio(S-2.5-dimethoxyphenyi)nicotinoyi-(*N*-2.5-dichlorobenzoyi)-*L*-4-aminophenyialanine

A solution of Intermediate 6 (0.44g, 0.89mmol) in THF (7.5ml) and water (5ml) was treated with lithium hydroxide monohydrate (43mg, 1.0mmol) and stirred at room temperature for 16h, then acidified to pH1 with 10% hydrochloric acid. The mixture was extracted with DCM (2 x 30ml) and the solvent evaporated in vacuo to give an off-white solid that was triturated with boiling MeOH to give the title compound as a white solid (210mg, 49%). &H (DMSO-d⁶) 11.18 (1H, br s, CO₂H), 10.86 (1H, s, NH), 8.89 (1H, d, J 8.0Hz), 8.30 (1H, dd, J 1.6, 4.7Hz), 7.77 (1H, dd, J 1.6, 7.6Hz), 6.96 (3H, m), 4.62 (1H, m), 3.70 (3H, s), 3.60 (3H, s) and 3.20-3.00 (2H, m), mvz (ES+) 626.628 (MH+).

The following compounds were prepared in a similar manner to the compound of Example 4a) by hydrolysis of the corresponding methyl ester. Each ester was obtained by coupling the starting materials shown according to the method described for intermediate 6:

a) 2-Thio(S-2,5-dimethoxyphenyl)nicotinoyl-(N-2,6-

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dichlorobenzoyl)-L-4-aminophenylalanine

from (N-2,6-dichlorobenzoy)-L-4-aminophenylalanine methyl ester and 3,5-dichloropyridyl-4-carbonyl chloride. δ H(DMSO-de) 8.64 (2H, s), 7.60-7.46 (5H, m), 7.24 (2H, d, \pm 8.5Hz), 4.78-4.65 (1H, m), 3.22-2.85 (2H, m); δ H m/z (ESI, 60V) 528 (MH+).

c) (<u>N'-2.4-Dimethylnicotinoyl)-(N-2.6-dichlorobenzoyl)-L-4-</u> aminophenylatenine

from (*N*-2.6-dichlorobenzoyl)-*L*-4-aminophenylalanine methyl ester and 2.4-dimethylpyridyl-4-carbonyl chloride. 8H (DMSO-d₆) 8.39 (1H, br d), 8.23 (1H, d, <u>J</u> 5.0Hz), 7.59-7.45 (6H, m), 7.26 (2H, d, <u>J</u> 8.4Hz), 7.01 (1H, d, <u>J</u> 5.1Hz), 4.65-4.52 (1H, m), 3.26-3.18 (1H, m), 2.95-2.84 (1H, m), 2.17 (3H, s); <u>ru</u>/z (ESI, 60V) 486 (MH+).

d) <u>N-(2,6-Dichloroisenicotinoyl)-C-(2,6-dichlorobenzyl)-L-tyrosine</u> from O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 2,6-dichloropyridyl-4-carbonyl chloride. &H (DMSO-d₆) 9.2 (1H, d), 7.81 (2H, s), 7.5-7.3 (3H,m), 7.21 (2H, d, <u>1</u> 8.5Hz), 6.93 (2H, d, <u>1</u> 8.5Hz), 5.15 (2H, s), 4.65 (1H, m), 3.28-3.15 (1H, m), 3.05-2.95 (1H, m); <u>m/z</u> (ESI, 60V)
 513 (MH+).

e) N'-(2-nicotinoyi)-O-(2,6-dichlorobenzyi)-L-tyrosine

from O-(2,6-dichlorobenzyl)-*L*-tyrosine methyl ester hydrochloride and nicotinoyl chloride using triethylamine as base. 8H (DMSO-d₆) 9.0-8.85 (2H, m), 8.7 (1H,m), 8.14 (1H, m), 7.55-7.41 (4H, m), 7.24 (2H, d, <u>J</u> 8.6Hz), 6.95 (2h, d, <u>J</u> 8.6Hz), 5.16 (2H, s), 4.59 (1H, m), 3.17-3.12 (1H, m), 3.04-2.96 (1H,m); m/z (ESI, 60V) 445 (MH+).

f) N-(3.5-Dichloro-4-picolyl)-N-(3,5-dichloro-4-picolyl)-L-4-amino-phenylalanine

from (N-3,5-dichloro-4-picolyl)-L-4-aminophenylalanine methyl ester and 3,5-dichlorophridyl-4-carbonyl chloride. 8H (DMSO-de, 300MHz) 9.26 (1H, d, <u>J</u> 8.3Hz), 8.79 (2H, s), 8.65 (2H, s), 7.57 (2H, d, <u>J</u> 8.4Hz), 7.30 (2H, d, <u>J</u> 8.4Hz), 4.70 (1H,m), 3.15 (1H, dd, <u>J</u> 14.1, 5.2Hz) and 2.93 (1H, dd, <u>J</u> 14.0, 9.3Hz); m/z (ESI, 160V) 527 (MH+).

g) <u>N-(2-Chloronicotinoyl)-N'-(3,5-dichloro-4-picolyl)-L-4-amino-phenylalanine</u>

from (N-3,5-dichloro-4-picolyl)-L-4-aminophenylalanine methyl ester and 2-chloro-nicotinoyl chloride. δH (DMSO-d₆, 300MHz)12.85 (1H, br s), 10.88 (1H,s), 8.97 (1H, d, $\frac{1}{2}$ 8.1Hz), 8.79 (2H, s), 8.46 (1H, dd, $\frac{1}{2}$ 4.8, 1.8Hz), 7.70 (1H, dd, $\frac{1}{2}$ 7.5, 1.8Hz), 7.59 (2H, d, $\frac{1}{2}$ 8.4Hz), 7.48 (1H, dd, $\frac{1}{2}$ 7.5, 4.8Hz), 7.30 (2H, d, $\frac{1}{2}$ 8.4Hz), 4.63 (1H, m), 3.16 (1H, dd, $\frac{1}{2}$ 13.9, 4.7Hz) and 2.95 (1H, dd, $\frac{1}{2}$ 13.8, 9.8Hz) m_Z (ESI, 160V) 493 (MH+).

10 EXAMPLE 5

a) Q-(2.6-dichlorobenzyl)-N-(4-acetyl-1,2,5-trimethyl-3-pyrroyl)-Ltyrosine

Intermediate 7 (360mg, 0.68mmol) was treated with LiQH.H₂O (34mg, 0.81mmol) in dioxane (6ml), water (6ml) and MeOH (4ml) at room temperature for 2h. The solvent was removed *in vacuo* and the obtained residue taken up in water. The pH was made acidic by addition of a few drops of acetic acid and the obtained precipitate filtered off with water washing affording the title compound as a white amorphous powder (245mg, 70%). &H (d⁶-DMSO) 8.37 (1H, d, ½ 8.2Hz, NH), 7.57-7.43 (3H, m's, aryl-H), 7.23 (2H, d, ½ 8.6Hz, aryl-H), 6.95 (2H, d, ½ 87.6Hz, aryl-H), 5.18 (2H, br s, CH₂-O), 4.62 (1H, m, xtyr-H), 3.33 (3H, s, MeN), 3.13 (1H, dd, ½ 4.1, 13.8Hz, CH₂H₂BAr), 2.32 (3H, s, MeCO), 2.01 (3H, s, pyrrole-Me) and 1.92 (3H, s, pyrrole-Me). m/z (ES+, 60V), 517 (MH+, 100), 519 (MH+, 70).

The following compounds were prepared in a similar manner to the compound of Example 5a) by hydrolysis of the corresponding methyl ester. Each ester was obtained by coupling the starting materials shown according to the method described for Intermediate 7:

b) <u>O-(2.6-dichlorobenzyl)-N-(4-acetyl-3,5-dimethyl-2-pyrroyl)-L-tyrosine</u>

from O-(2,6-dichlorobenzyl)-*L*-tyrosine methyl ester hydrochloride and 3,5-dimethyl-4-acetylpyrrole-2-carboxylic acid. Freeze drying afforded the <u>title compound</u> as a light cream amorphous solid (550mg). 8H (DMSO-d₆) 11.56, (1H, s), 7.61 (1H, d, <u>J</u> 7.8Hz), 7.51 (2H, d, <u>J</u> 8.0Hz), 7.41 (1H, t, <u>J</u>

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8.0Hz), 7.21 (2H, d, J 85Hz), 6.96 (2H, d, J 8.5Hz), 5.16 (2H, s), 4.69-4.55 (1H, m), 3.12 (1H, dd, J 13.7, 4.7Hz), 2.99 (1H, dd, J 13.7, 9.1Hz), 2.43 (3H.s), 2.38 (3H, s), 2.31 (3H, s); m/z (ESI, 60V) 503 (MH+).

5 c) O-(2.6-dichlorobenzyl)-N-(4-acetyl-2,5-dimethyl-3-pyrrolyl)-Ltyrosine

from O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 2,5dimethyl-4-acetyl-pyrrole-3-carboxylic acid. Feeze drying afforded the title compound as a white amorphous solid (203mg), δH (DMSO-de) 11.2 (1H. s), 8.83 (1H, d, <u>J</u> 8.0Hz), 7.56 (2H, app.d. <u>J</u> 8.0Hz), 7.45 (1H, app.t, <u>J</u> 8.0Hz), 7.21 (2H, d, J 8.5Hz), 6.95 (2H, d, J 8.5Hz), 5.17 (2H, s), 4.61-4.52 (1H, m), 3.11 (1H, dd, J 13.8, 4.3Hz), 2.86 (1H, dd, J 13.8, 10.4Hz), 2.31 (3H, s), 2.08 (3H, s), 2.07 (3H, s); m/z (ESI, 60V) 503 and 505 (MH+).

15 O-(2.6-dichlorobenzyl)-N-(1-methyl-2-indolyl)-L-tyrosine from O-(2.6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 1methylindole-2-carboxylic acid. Freeze drying afforded the title compound as a white amorphous solid (220mg). δH (DMSO-d₆) 11.7 (1H, br s), 8.66 (1H, d, J 8.3Hz), 7.65 (1H, d, J 7.9Hz), 7.53 (2H, app.d, J 8l1Hz), 7.26 20 (1H,obscured m), 7.09 (2H, app.t, J 7.5Hz), 6.96 (2H, d, J 8.5Hz), 5.16

(2H, s), 4.64-4.54 (1H, m), 3.89 (3H, s), 3.16 (1H, dd, J 13.8, 4.3Hz) and 3.00 (1H, dd, J 13.8, 10.4Hz); m/z (ESI, 60V) 497 and 499 (MH+).

O-(2.6-dichlorobenzyl)-N-(2-(4-chlorophenyl)-3-(trifluoromethyl) e) -4-pyrazovii-L-tyrosine

from O-(2.6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 2-(4-chlorophenyl)-3-(trifluoromethyl)-pyrazole-4-carboxylic acid. The title compound was isolated as an off-white solid (220mg). δH (DMSO-d₆) 11.17 (1H, br s), 8.82 (1H, d, J 8.2Hz), 8.08 (1H, s), 7.65 (2H, d, J 8.1Hz), 7.54 (4H, app. d. J 8.1Hz), 7.45 (1H, app. t. J 8.0Hz), 7.24 (2H, d, J 8.3Hz), 6.98 (2H, d, J 8.3Hz), 5.18 (2H, s), 4.61-4.51 (1H, m), 3.17 (1H, dd, <u>J</u> 13.8, 4.5Hz) and 2.94 (1H, dd, <u>J</u> 13.8, 9.0Hz); <u>m/z</u> (ESI, 60V) 527 and 529 (MH+).

35 O-(2.6-dichlorobenzyl)-N-(2-phenyl-4-thiazovl)-L-tyrosine f)

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from O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 2-phenyl-thiazole-4-carboxylic acid. Freeze-drying afforded the title compound as a pale yellow amorphous solid (340mg). δ H (DMSO-dg) 11.2 (1H, br s), 8.40 (1H, d, \downarrow 8.1Hz), 8.31 (1H, s) 8.05-8.02 (2H, m), 7.55-7.51 (5H, m), 7.46-7.40 (1H, m), 7.23 (2H, d, \downarrow 8.0Hz), 6.97 (2H, d, \downarrow 7.8Hz), 5.16 (2H, s), 4.71-4.64 (1H, m), 3.18 (2H, app. d, \downarrow 6.6Hz); m/z (ESI, 60V) 527 and 529 (MH*).

g) (N'-1-Methyl-5-nitropyrazolyl)-(N-2.6-dichlorobenzoyl)-L-4aminophenyalanine

from (*N*-2,6-dichlorobenzoyl)-*L*-4-aminophenylalanine methyl ester with *N*-methyl-5-nitropyrazole-4-carboxylic acid. δ H (DMSO-d₆) 10.64-(1H, s), 8.68 (1H, br d), 7.80 91H, s), 7.64-7.48 (6H, m), 7.19 (2H, d, \pm 8.5Hz), 4.59-4.49 (1H, m), 4.05 (3H, s), 3.18-2.9 (2H, m); $\underline{m/z}$ (ESI, 60V) 506 (MH+).

- h) M-(2-Methylnicotinovi)-O-(2,6-dichlorobenzyi) L-tyrosine from O-(2,6-dichlorobenzyi)-L-tyrosine methyl ester hydrochloride and 2-methylnicotinic acid. &H (DMSO-de) 8.73 (1H, br d), 8.48 (1H, m), 7.7.46 (4H, m), 7.4-7.23 (3H, m), 7.00 (2H, d, 1 8.4Hz), 5.2) (2H, s), 4.60 91H, m), 3,27-3.12 (1H, m), 3,0-2.82 (1H, m); m/z (ESI, 60V) 509 (MH+).
- i) (N*2-Chloronicotinoyl)-(N*benzoyl)-L-4-sminophenylalanine from (N*-benzoyl)-L-4-aminophenylalanine methyl ester and 2-chloronicotinic acid. 8H (DMSO-de) 10.19 (1H, s), 8.96 (1H, d, J 8.2Hz), 8.46 (1H, dd, J 4.8, 1.9Hz), 7.96 (2H, dd, J 6.7, 1.7H), 7.75-7.65 (3H, m), 7.62-7.45 (4H, m), 7.25 (2H, d, J 8.5Hz), 4.6 (1H, m), 3.2-3.12 (1H, m), 3.0-2.89 (1H, m), m/z (ESI, 60V) 424 (MH+).
- j) N'-(Quinoline-4-carbonyl)-C-(2.6-dichlorobenzyl)-L-tyrosine from O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 4-quinoline carboxylic acid. 8H (DMSO-de) 9.0 (1H, d), 8.94 (1H, d, <u>1</u> 4.3), 8.06 (1H, d, <u>1</u> 8.2Hz), 7.79 (2H, m), 7.62-7.42 (4H, m), 7.34 (1H, d, <u>1</u> 4.3Hz), 7.34 (2H, d, <u>1</u> 8.7Hz), 7.00 (2H, d, <u>1</u> 8.7Hz), 5.22 (2H, s), 4.72 (1H, 35 m), 3.29-3.19 (1H, m), 3.0-2.98 (1H, m); m/z (ESI, 60V) 495 (MH+).

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- k) <u>A'-(2-Phenoxynicotinoyl)-O-(2,6-dichlorobenzyl)-L-tyrosine</u> from O-(2,6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 2-phenoxynicotinic acid. δH (DMSO-de) 8.55 (1H, d), 8.18 (2H, m), 7.6-7.4 (5H, m), 7.3-7.15 (6H,m), 6.71 (2H, d), 5.1 (2H, s), 4.65 (1H, m), 3.22-3.0 (2H, m): m/z (ESI, 60V) 537 (MH+).
- M-(Pyridine-2-carbonyl)-O-(2.6-dichlorobenzyl)-L-tyrosine from O-(2.6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and 2-picolinic acid. 8H (DMSO-dg) 8.8-8.6 (2H, m), 8.0 (2H, m), 7.7-7.4 (4H, m),
 7.14 (2H, d, <u>J</u> 8.7Hz), 6.92 (2H, d, <u>J</u> 8.7Hz), 5.15 (2H, s), 4.72 (1H, m),
 3.17 (2H, m); <u>m/z</u> (ESI, 60V) 445 (MH+).
 - m) N-(Pyridine-4-carbonyl)-O-(2.6-dichlorobenzyl)-L-tyrosine from O-(2.6-dichlorobenzyl)-L-tyrosine methyl ester hydrochloride and isonicotinic acid. δH (DMSO-d₆) 8.68 (2H, dd, <u>J</u> 4.5, 1.6Hz), 8.4 (1H, d, <u>J</u> 7.2Hz), 7.63 (2H, dd, <u>J</u> 4.5, 1.6Hz), 7.6-7.4 (3H, m), 7.15 (2H, d, <u>J</u> 8.6Hz), 6.87 (2H, d, <u>J</u> 8.6Hz), 5.14 (2H, s), 4.31 (1H, m), 3.2-2.96 (2H, m); <u>m/z</u> (ESI, 60V) 445 (MH+).
 - n) <u>M-(2-Hydroxynicotinoyl)-*D*-(2,6-dichlorobenzyl)-*L*-tyrosine)</u> from O-(2,6-dichlorobenzyl)-*L*-tyrosine methyl ester hydrochloride and 2-hydroxynicotinic acid. δ H (DMSO-d₆) 10.15 (1H, d, \pm 7.4Hz), 8.31 (1H, dd, \pm 7.1, 2.0Hz), 7.71 (1H, br s), 7.65-7.45 (3H, m), 7.16 (2H, d, \pm 8.4Hz), 6.97 (2H, d, \pm 8.4Hz), 6.46 (1H, t, \pm 6.8Hz), 5.17 (2H, s), 4.68 (1H, m), 3.2-3.0 (2H, m); <u>m/z</u> (ESI, 60V) 461 (MH+).
 - o) (N-2-Aminonicotlnoyl)-(N-2,6-dichlorobenzoyl)-L-4-amino phenylalanine

from (N-2,6-dichlorobenzoyl)-L-4-aminophenylalanine methyl ester and 2-aminonicotinic acid. 8H (DMSO-d₆) 10.64 (1H, s), 8.55 (1H, d), 8.06 (1H, m), 7.8 (1H, d), 7.7-7.4 (4H, m), 7.28 (2H, d, 过 8.5Hz), 6.9 (2H, br s), 6.5 (1H, m), 4.52 (1H, m), 3.2-2.9 (2H, m); m/z (ESI, 60V) 473 (MH+).

p) (N-2-Hydroxynicotinoyl)-(N-3.5-dichloroisonicotinoyl)-L-4aminophenylalanine

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from (*N*-3,5-dichloroisonicotinoyl)-*L*-4-aminophenylalanine methyl ester and 2-hydroxynicotinic acid. 8H (DMSO-d₆) 10.14 (1H, d, <u>J</u> 7.6Hz), 8.78 (2H, s), 8.31 (1H, dd, <u>J</u> 7.2, 2.2Hz), 7.70 (1H, br m), 7.57 (2H, d, <u>J</u> 8.5Hz), 7.20 (2H, d, <u>J</u> 8.5Hz), 6.46 (1H, m), 4.69 (1H, m), 3.20-2.95 (2H, m); <u>m/z</u> (ESI, 60V) 475 (MH+).

q) (N'-2-Methylnicotinoyl)-(N-3,5-dichloroisonicotinoyl)-L-4aminophenylalanine

from (Ar-3,5-dichloroisonicotinoy)-L-4-aminophenylalanine and 2-methylnicotinic acid. 8H (DMSO-d₆) 8.78 (2H, s), 8.60 (1H, br d), 8.48 (1H, dd, <u>1</u> 4.8, 1.6Hz), 7.7-7.5 (3H, m), 7.4-7.2 (3H, m), 4.5 (1H,), 3.3-2.85 (2H, m), 2.34 (3H, s); m/z (ESI, 60V) 473 (MH+).

r) (N-2,5-Dichlorobenzoyi)-(N-2-phenoxynicotinoyi)-L-4aminophenyialanine

from (N-2,6-dichlorobenzoyi)-L-4-aminophenylalanine methyl ester and 2phenoxynicotinic acid. δH (DMSO-d₀) 10.61 (1H, s), 8.56 (1H, d, ป 7.5Hz), 8.22-8.17 (2H, m), 7.59-7.39 (8H, m), 7.27-7.11 (7H, m), 4.72-4.64 (1H, m, CH), 3.16 (1H, dd, ป 13.7,4.9Hz, CHAH_B) and 3.06 (1H, dd, ป 13.7, 7.8Hz,CHAH_B); m/z (ESI, 60V) 550 (MH*).

s) (N3.5-Dichloroisonicotinoyl)-(N2-phenoxynicotinoyl)-L-4-amionophenylalanine

from (N-3,5-dichlorolsonicotinoyl)-L-4-aminophenylalanine methyl ester and 2-phenoxynicotinic acid. ∂H (DMSO-de) 10.79 (1H, s), 8.78 (2H, s, ArH), 8.57 (1H, d, ⊥ 7.6Hz), 8.25-8.13 (2H, m), 7.46-7.35 (4H, m), 7.29-7.05 (6H, m), 4.73-4.82 (1H, m, CH), 3.17 (1H, dd, ⊥ 13.7, 4.7Hz, CHAHs) and 3.09 (1H, dd, ⊥ 13.7, 7.8Hz, CHAHs); m/z (ESI, 60V) 551 (MH+*).

30 t) N-(4-Acatyl-1,2,5-trimethyl-1*H*-pyrrole-3-carbonyl)-*N*-(3,5-dichloro-4-picolyl)-*L*-4-aminophenylalanine

from (N-3,5-dichloro-4-picolyf)-L-4-aminophenylalanine methyl ester and 4acetyl-1.2,5-trimethyl-1*H*-pyrrole-3-carboxylic acid. δH (DMSO-de, 300MHz) 400K) 12.72 (1H, br s), 10.83 (1H,s), 8.80 (2H, s), 8.42 (1H, d, <u>J</u> 8.4Hz), 7.54 (2H, d, <u>J</u> 8.5Hz), 7.30 (2H, d, <u>J</u> 8.4Hz), 4.66 (1H, m), 3.34 (3H, s), 3.16 (1H, dd, <u>J</u> 13.8, 4.3Hz), 2.89 (1H, dd, <u>J</u> 13.8, 11.0Hz), 2.32 (3H, s), 2.05 (3H, s) and 1.90 (3H, s); m/z (ESI, 60V) 531 (MH+).

u) <u>N-(4-(Carboxy)nicotinoyl)-N'-(3,5-dichloro-4-picolyl)-L-4-</u> aminophenylalanine

from (N-3,5-dichloro-4-picolyl)-L-4-aminophenylalanine methyl ester and 4-(methoxycarbonyl)nicotinic acid . \$H (DMSO-d₆, 300MHz) 10.98 (1H, s), 9.01 (1H, d, \pm 8.0Hz), 8.77 (2H,s), 8.63 (1H, s), 7.66 (1H, d, \pm 5.1Hz), 7.59 (2H, d, \pm 8.5Hz), 7.32 (2H, d, \pm 8.5Hz), 4.63 (1H, m), 3.14 (1H, dd, \pm 13.9, 5.4Hz) and 3.03 (1H, dd, \pm 13.9, 8.8Hz); m/z (ESI, 160V) 503 (MH+).

v) (2-Acetyl-3-thienyl)carbonyl-(N-3,5-dichloro-4-picollnyl)-4-aminophenylalanine

from (N-3,5-dichloro-4-picolyl)-L-4-aminophenylalanine methyl ester and 2-acetyl-thiophene-3-carboxylic acid. δH (DMSO-de) 9.17 (1H, d, \downarrow 8.2Hz, NH), 8.77 (2H, s, pyr-H), 7.99 (1H, d, \downarrow 5.1Hz, thiophene H-5), 7.57 (2H, ABd, \downarrow 8.5Hz, Ar-H), 7.31 (2H, ABd, 2H, \downarrow 8.5Hz, Ar-H), 7.07 (1H, d, \downarrow 5.1Hz, thiophene H-4), 5.38 (1H, m, CH α) 3.18 (1H, dd, \downarrow 13.8, 4.6Hz, CHCHAHBAr), 2.93 (1H, dd, \downarrow 13.8, 10.3Hz, CHCHAHBAr) and 2.26 (3H, s, COMe); m/z (ESI, 60V) 506 (MH+).

EXAMPLE 6

2-Chloronicotinoyl-(N-2,6-dichlorobenzoyl)-L-4aminophenylalanine

Lithium hydroxide monohydrate (109mg, 2.5mmol) was added to a solution of Intermediate 8 (450mg, 1mmol) in a mixture of THF (10ml) and water (10ml). The mixture was stirred for 2h at room temperature, then the THF was evaporated in vacuo. The aqueous residue was neutralised (11M hydrochloric acid), and the precipitate isolated by filtration, washed with water and dried to give the title compound (300mg, 61%). &H (DMSO-d6, 400K) 8.90 (1H, d, \downarrow 9.1Hzm 1NH), 8.48 (1H, m, Py H), 7.69 (1H, m, Py H), 7.64-7.45 (7H, m, 4ArH, 1 NH, 1 PyH), 7.29 (2H, d, \downarrow 8.3Hz, 2 ArH), 4.66-4.53 (1H, m, Chatyr), 3.18 (1H, dd, \downarrow 14, 5.2Hz CHAHBAr), and2.91 (1H, dd, \downarrow 9.6, 14Hz, CHAHBAr), m/z (ESI, 60V) 491 (MH+).

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The following compounds were prepared in a similar manner:

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b) M-(2-Chloronicotinoyl)-O-(2.6-dichlorobenzoyl)-L-tyrosine from Intermediate 30 to give the title compound as a white solid. δH (DMSO-de, 300MHz) 12.91 (1H, br s, GO₂H), 8.98 (1H, d, H 8.3Hz, CONH), 8.44 (1H, dd, J 4.8, 1.9Hz, PyH), 7.69-7.58 (4H, m, Cl₂AH₃ + PyH), 7.45 (1H, dd, J 7.5, 4.9Hz, PyH), 7.43 (2H, d, J 8.7Hz, ArH), 7.22 (2H, d, J 8.5Hz, ArH), 4.67 (1H, ddd, J 10.0, 8.2, 4.7Hz, CHα), 3.24 (1H, dd, J 14.0, 4.0Hz, CH_AH_BAr) and 2.99 (1H, dd, J 13.9, 10.2Hz, CH_AH_BAr); m/z (ESI, 60V) 493 (MH+).

N-(2-Chioronicotinoyl)-N-methyl-N'-(3.5-dichloro-4-picolyl)-L-4aminophenylalanine

from Intermediate 31 to give the <u>title_compound</u> as a white solid δH (DMSO-d₆, 300MHz, 405K) 10.36 (1H, br s, CONH), 8.67 (2H, s, Cl₂PyH), 8.42–8.39 (1H, m, ClPyH), 7.54-7.15 (6H, br m, 4 x ArH + 2 x ClPyH), 5.30 (1H, v br s, CH α), 3.4-2.6 (5H, br m, NMe + CHC \underline{H}_2 Ar) (Acid proton not observed at 405K, at 300K δH 13.06 (1H, br s, CO₂H)): m/z (ESI, 70V) 507 (MH+).

d) [(S-2,5-dimethoxyphenyl)sulphonyl]nicotinoyl-O-(2,5-dichlorobenzyl)-L-tyrosine

e) 2-(((2-Chloro-3-pyridinyl)carbonyl]amino)-3-(4-((2,6-

dichloroanilino)carbonyliphenyl)propanoic acid from Intermediate 15 to give the <u>title compound</u> as an off white solid δH (DMSO-d₆, 300K) 11.06 (1H, br s), 10.23 (1H, br s), 9.00 (1H, d, \downarrow 7.8Hz), 8.46 (1H, br d), 7.95 (2H, d, \downarrow 7.4Hz), 7.69 (1H, d, \downarrow 7.1Hz), 7.59 (2H, d, \downarrow 8.0Hz), 7.50-7.36 (3H, m), 4.71 (1H, br), 3.26 (1H) and 3.06 (1H, dd, \downarrow 13.8, 10.0Hz): m/z (ESI, 60V) 492 (MH*).

f) 2-[[(2-Chloro-3-pyridinyl)carbonyl]amino}-3-(4-{[(3,5-dichloro-4-pyridinyl)amino]carbonyl]phenyl propionic acid

from the corresponding inermediate ester prepared in a similar way to Intermediate 15 to give <u>title compounds</u> as an offwhite solid. δH (DMSOde, 300MHz) 12.92 (1H, br s), 10.57 (1H, s), 9.00 (1H, d, \downarrow 8.2Hz), 8.75 (2H, s), 8.45 (1H, dd, \downarrow 4.7, 1.7Hz), 7.96 (2H, d, \downarrow 6.15Hz), 7.69 (1H, d, \downarrow 7.3, 1.7Hz), 7.49 (2H, d, \downarrow 8.0Hz), 7.48 (1H, 4.71 (1H, br m), 3.28 (1H, dd, \downarrow 13.9, 4.7Hz) and 3.06 (1H, dd, \downarrow 13.9, 10.0Hz); <u>m/z</u> (ESI, 60V) 483 (MH+).

EXAMPLE 7

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2-Thio(S-acetic acid)nicotinovi-O-(2.6-dichlorobenzyi)-L-tyrosine

Lithium hydroxide monohydrate (75mg, 1.8mmol) was added to a solution of Intermediate 9 (360mg, 0.6mmol) in a mixture of THF (13ml) and water (10ml). The mixture was stirred for 2hr at room temperature, then the THF was evaporated *in vacuo*. The aqueous residue was neutralised (1M hyrochloric acid), and the precipitate isolated by filtration, washed with water and dried to give the <u>title compound</u> (200mg, 58%). δ H (DMSO-d₆), 400K), 8.5-8.35 (2H, m, pyrH, 1NH), 7.71 (1H, dd, \downarrow 1.7, 7.6, pyrH), 7.57 (2H, d, \downarrow 8.9, 2ArH), 7.45 (1H, m, 1ArH), 7.22 (2H, d, \downarrow 8.5, 2ArH), 7.18 (1H, m, 1pyrH), 6.97 (2H, d, \downarrow 8.5, 2ArH), 5.18 (2H, s, OCH₂Ar), 4.43 (1H, m, CHatyr), 3.82 (2H, s, SCH₂CO₂H), 2.94-3.23 (2H, m, CH₂Ar); m/z (ESI, GOU) 535 (MH+).

25 EXAMPLE 8

2-Thio(S-methyl)nicotinovI-O-(2.6-dichlorobenzyl)-L-tyrosine

Lithium hydroxide monohydrate (140mg, 3.3mmol) was added to a solution of Intermediate 10 (1.4gm, 2.7mmol) in a mixture of THF (10ml) and water (10ml). The mixture was stirred for 2 hr at room temperature then the THF was evaporated in vacuo. The aqueous residue was neutralised (1M hydrochloric acid), and the precipitate isolated by filtration, washed with water and dried to give the <u>title compound</u> (1.1gm, 81%). δH (DMSO-de, 400K) 8.73 (1H, d, <u>J</u> 8.1, NH), 8.52 (1H, dd, <u>J</u> 1.7, 4.8, 1pyrH), 7.67 (1H, dd, <u>J</u> 1.7, 7.6, 1pyrH), 7.55 (2H, d, <u>J</u> 8.9, 2ArH), 7.45 (1H, dd, <u>J</u> 2.3, 8.9, 1ArH), 7.24 (2H, d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, <u>J</u> 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99 (2H,d, J 8.6, 2ArH), 7.16 (1H, m, 1pyrH), 6.99

2ArH), 5.18 (2H, s, OCH₂Ar), 4.55 (1H, m, CH₂tyr), 3.16-2.95 (2H, m, CH₂Ar) and 2.38 (3H, s, SCH₃); m/z (ESI, GOU) 491 (MH+).

EXAMPLE 9

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(N-2-Chloronicotinoyi)-4-[(E)-2-(2,6-dichlorophenyi)ethenyi]

To a solution of Intermediate 28 (1.0g, 1.99mmol) in THF (5ml) and water (5ml) was added lithium hydroxide monohydrate (88mg 2.09mmol). The reaction mixture was stirred for 1h. The THF was then removed *in vacuo* and the remaining aqueous solution acidified to pH6 with 1M hydrochloric acid. The resulting precipitate was collected and washed with water and ether and finally freeze dried. The resulting compound contained an impurity so a small amount was purified by preparavie HPLC (98mg). δ H (DMSO-d₆) 12.90 (1H, dr s, CO₂H), 8.98 (1H, d, \downarrow 8.0Hz), 8.45 (1H, d, \downarrow 3.4Hz), 7.70-7.03 (9H, m), 4.72-4.60 (1H, m, CH), 3.20 (1H, dd, \downarrow 14.0, 4.5Hz, CH_AH_B) and 3.01 (1H, dd, \downarrow 14.0, 9.9Hz, CH_AH_B); m/z (ESI, 60V) 475 (MH+).

EXAMPLE 10

20 (N-2-Chloronicotinoyl)-4-[2-(2,6-dichlorophenyl)-2-hydroxyethyl] phenylalanine

Lithium hydroxide monohydrate (13mg) was added to a solution of Intermediate 22 (150mg, 0.29mmol) in THF (5ml) and H₂O (5ml). The solution was stirred for 1h and then the THF removed *in vacuo* and the remaining aqueous solution acidified to pH6 with 1M hydrochloric acid. The solid precipitate formed was collected by filtration, washed with copious quantities of water and finally freeze dried to give the title compund as a fluffy white solid (70mg, 49%). &H (DMSO-da) 12.78 (1H, br s, CO₂H), 8.90 (1H, d, J 8.0Hz), 8.45 (1H, d, J 4.8Hz), 7.63-7.58 (1H, m, ArH), 7.49-7.45 (1H, m, ArH), 7.34-7.07 (7H, m, ArH), 5.50-5.45 (1H, m, CH), 4.61-4.51 (1H, m, CH) and 3.30-2.35 (4H, m, 2 x CH₂); m/z (ESI, 60V) 493 (MH+).

EXAMPLE 11

35 (N-2-Chloronicotinoyl)-[4-[2-(2-dichlorophenyl)-2-excethyl]): phenylalanine

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Lithium hydroxide monohydrate (36mg, 0.85mmol) was added to a solution of Intermediate 23 (400mg, 0.77mmol) in THF (5ml) and water (5ml). The reaction mixture was stirred for 3h and then the THF was removed *invacuo*. The remaining aqueous solution was acidified with 1M hydrochloric acid. The resulting white precipitate was collected and washed well with water. Further purification by column chromatography (SiO₂; acetic acid: MeOH:DCM,2:8:90) gave the <u>title compound</u> as a white solid (78mg, 19%). &H (DMSO-d₆) 8.45 (1H,dd, <u>1</u>4.8, 2.0Hz), 8.03 (1H,dd, <u>1</u>7.7, 2.0Hz), 7.36-7.20 (8H, m, ArH), 6.98 (1H, d, <u>1</u>7.2Hz, ArH), 5.14-5.05 (1H, m, CH), 4.11 (2H, s. CH₂C=O)m, 3.39 (1H, dd, <u>1</u>14.1, 5.6Hz, CH₂H₈) and 3.25 (1H, dd, <u>1</u>14.1, 6.2Hz, CH₂H₈); m/z (ESI, 60V) 491 (MH*).

α4β1 Integrin-dependent Jurkat cell adhesion to VCAM-ig

96 well NUNC plates were coated with F(ab)₂ fragment goat anti-human IgG Fcγ-specific antibody [Jackson Immuno Research 109-006-098: 100 μl at 2 μg/ml in 0.1M NaHCO₃, pH 8.4], overnight at 4°. The plates were washed (3x) in phosphate-buffered saline (PBS) and then blocked for 1h in PBS/1% BSA at room temperature on a rocking platform. After washing (3x in PBS) 9 ng/ml of purified 2d VCAM-Ig diluted in PBS/1% BSA was added and the plates left for 60 minutes at room temperature on a rocking platform. The plates were washed (3x in PBS) and the assay then performed at 37° for 30 min in a total volume of 200 μl containing 2.5 x 105 Jurkat cells in the presence or absence of titrated test compounds.

Each plate was washed (2x) with medium and the adherent cells were fixed with 100µl methanol for 10 minutes followed by another wash. 100µl 0.25% Rose Bengal (Sigma R4507) in PBS was added for 5 minutes at room temperature and the plates washed (3x) in PBS. 100µl 50% (v/v) ethanol in PBS was added and the plates left for 60min after which the absorbance (570mm) was measured.

αμβη Integrin-dependent JY cell adhesion to MAdCAM-ig

This assay was performed in the same manner as the α4β1 assay except
that MAdCAM-Ig (150ng/ml) was used in place of 2d VCAM-Ig and a subline of the β-lympho blastoid cell-line JY was used in place of Jurkat cells.

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The IC50 value for each test compound was determined as described in the $\alpha_A \beta_1$ integrin assay.

α₅β₁ Integrin-dependent K562 cell adhesion to fibronectin

96 well tissue culture plates were coated with human plasma fibronectin (Sigma F0895) at $5\mu g/ml$ in phosphate-buffered saline (PBS) for 2 hr at 37°C. The plates were washed (3x in PBS) and then blocked for 1h in 100 μ i PBS/1% BSA at room temperature on a rocking platform. The blocked plates were washed (3x in PBS) and the assay then performed at 37°C in a total volume of 200 μ l containing 2.5x 10⁵ K562 cells, phorbol-12-myristate-13-acetate at 10ng/ml, and in the presence or absence of titrated test compounds. Incubation time was 30 minutes. Each plate was fixed and stained as described in the $\alpha 4\beta$ 1 assay above.

α_mβ₂-dependent human polymorphonuclear neutrophils adhesion to

96 well tissue culture plates were coated with RPMI 1640/10% FCS for 2h at 37°C. 2 x 10°5 freshly isolated human venous polymorphonuclear neutrophils (PMN) were added to the wells in a total volume of 200µl in the presence of 10ng/ml phorbol-12-myristate-13-acetate, and in the presence or absence of test compounds, and incubated for 20min at 37°C followed by 30min at room temperature. The plates were washed in medium and 100µl 0.1% (w/v) HMB (hexadecyl trimethyl ammonium bromide, Sigma H5882) in 0.05M potassium phosphate buffer, pH 6.0 added to each well. The plates were then left on a rocker at room temperature for 60 min. Endogenous peroxidase activity was then assessed using tetramethyl benzidine (TMB) as follows: PMN lysate samples mixed with 0.22% H2O2 (Sigma) and 50μ g/ml TMB (Boehringer Mannheim) in 0.1M sodium acetate/citrate buffer, pH 6.0 and absorbance measured at 630nm.

allb/82 -dependent human platelet aggregation

Human platelet aggregation was assessed using impedance aggregation on the Chronolog Whole Blood Lumiaggregometer. Human platelet-rich plasma (PRP) was obtained by spinning fresh human venous blood anticoagulated with 0.38% (v/v) tri-sodium citrate at 220xg for 10 min and diluted to a cell density of 6 x 109/ml in autologous plasma. Cuvettes

contained equal volumes of PRP and filtered Tyrode's buffer (g/liter: NaCl 8.0; MgCl $_2$ -H $_2$ O 0.427; CaCl $_2$ 0.2; KCl 0.2; D-glucose 1.0; NaHCO $_3$ 1.0; NaHPO $_4$ -2H $_2$ O 0.065). Aggregation was monitored following addition of 2.5 μ M ADP (Sigma) in the presence or absence of inhibitors.

In the above assays the compounds of the invention generally have IC_{50} values in the $\alpha_4\beta_1$ and $\alpha_4\beta_7$ assays of 1 μ M and below. Thus compounds of the Examples typically had IC_{50} values of 100nM and below in these assays and demonstrated selective inhibition of $\alpha4\beta_1$. In the other assays featuring α integrins of other subgroups the same compounds had IC_{50} values of 50 μ M and above thus demonstrating the potency and selectivity

of their action against ou integrins.